CONGENITAL DIABETES MELLITUS AND NEONATAL PSEUDODIABETES MELLITUS

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Diabetes mellitus is very rare in infants. Cases in recent years have been reported by Schwartzman, Crusius and Beirne (1947), by Bertoye and Stefani (1954) and by Keidan (1955), who reviewed the hitherto published cases of infantile diabetes. In Keidan’s report 79 cases of infantile diabetes are mentioned, the neonatal cases attracting special interest on account of certain clinical characteristics.

These cases have also been published under such varying headings as congenital diabetes (Ambard, Merklen, Schmid, Wolf and Arnowievitch, 1925; Feldmann, 1928; Nevinny and Schretter, 1930; Schretter and Nevinny, 1930; Wylie, 1953) and neonatal diabetes mellitus (Lewis and Eisenberg, 1935). Several of the congenital cases have been but insufficiently investigated, and Lawrence and McCance (1931) by an exact clinical analysis could reject a number of these as being unproved congenital cases. In most cases falling in this group (Kitselle, 1852; Morrison, 1916; Ambard et al., 1925; Feldmann, 1928; Schretter and Nevinny, 1930; Nevinny and Schretter, 1930) the diabetic mother as well as the child died at parturition.

Of proved cases of congenital diabetes there is the case reported by Guest (1949) in which diabetes was diagnosed on the ninth day after parturition, and Wylie’s (1953) case discovered at the age of 2 weeks. Both of these cases resulted in permanent diabetes. The remaining cases of neonatal diabetes may be divided into two groups, the first of which may be called neonatal diabetes mellitus and into which can be put cases reported by Cuno (1910), Kochmann (1922) (these two cases are, however, doubtful) Lewis and Eisenberg (1935), Limper and Miller (1935), Devine (1938), Gans (1954), Hofman-Bang (1954) and Hickish (1956). Most of these cases of diabetes have had an abrupt course with a fatal outcome. The other group comprises diseases which differ not only from the other cases of neonatal diabetes, but also from all other instances of diabetes, because they all had a good prognosis and all were cured (Ramsey, 1926; Lawrence and McCance, 1931; Strandquist, 1932; Nawrocka-Kańska, 1952; Arey, 1953; Keidan, 1955; and Nelson, 1954). The majority simultaneously showed a marked infectious state which, however, was not considered of primary aetiological importance. The abnormal carbohydrate metabolism has generally lasted two or three weeks only and as a rule has yielded to insulin therapy. In a few cases a cerebrally conditioned diabetic state has been suspected (Nawrocka-Kańska, 1952; Keidan, 1955).

They have been described under various designations, for example, ‘temporary diabetes’ (Lawrence and McCance, 1931), ‘transient diabetes’ (Arey, 1953; Keidan, 1955). Strandquist (1932) speaks of ‘infantile glycosuria simulating diabetes’ and Nelson (1954) of ‘diabetes mellitus syndrome in the newborn infant’. As a common name for this condition we suggest ‘neonatal pseudodiabetes mellitus’.

Case Reports

Case 1. A boy was born on February 3, 1953, at the Obstetric Department in Lund. The parents were healthy, without any diabetic heredity. The pregnancy had a normal course, but the patient was delivered by means of low forceps on account of threatening foetal hypoxia. His birth weight was 2,780 g. (6 lb. 2 oz.) and length at birth 49 cm. (19 in.). The child showed pronounced signs of postmaturity (Fig. 1) and was referred to stage III
in Clifford's (1954) classification of placental dys-
function.

The patient's weight fell to 2,300 g. (5 lb.) and he 
became cachectic, although alert. He began to vomit 
and on the fifth day after parturition was transferred to 
the Pediatric Department in Lund suspected of an 
endocrine-metabolic disorder. Glycosuria was proved, 
non-protein nitrogen was 35 mg./100 ml. and the 
blood sugar was as high as 720 mg./100 ml. In the course 
of the following days the blood sugar was about 400 to 
500 mg./100 ml., the glycosuria was maximally 13\% but 
there was no ketonuria.

At first the patient was fed on small quantities of breast 
milk, with plenty of fluid and electrolytes. Testosterone 
was also administered. On this therapy the weight was 
maintained unchanged, the blood sugar was about 
500 mg./100 ml. but the patient showed no ketosis and 
the general condition was surprisingly good. More 
breast milk was now allowed and the blood sugar was 
further increased to about 600 mg./100 ml. and at the 
same time Legal's test became positive.

At 3 weeks of age insulin therapy was instituted.

Initial laboratory tests showed eosinopenia, as well as 
signs of hypokalaemia. Cholesterol and phospholipids 
were pathologically increased, 254 and 454 mg./100 ml. 
respectively. The sugar in the cerebrospinal fluid was 
380 mg./100 ml. Insulin, starting with 4 i.u. NPH-insulin 
was given, the dose being gradually increased to 8 i.u. 
which was soon reduced again to 6 and finally to 4 i.u. 
On the second day the patient was ketone-free and the 
bleed sugar values fell to between 250 and 300 mg./100 ml. 
After three weeks we changed over to the same dosage 
of 4 i.u. IZS (Lente) and after the lapse of a little less 
than a week the patient showed no sugar in the urine 
and the blood sugar was normal.

Two weeks later we experimentally omitted the insulin 
for four days. Immediately there was glycosuria, and 
the blood sugar rose abruptly to 350 to 400 mg./100 ml. 
and for this reason we again began to administer insulin 
in the same dosage, 4 i.u., first NPH and later, after two 
weeks, IZS (Lente). Some days after the change-over 
to IZS (Lente) the blood sugar was again normal and 
the urine free of sugar. The insulin was continued for 
another three weeks and then was discontinued for good. 
At that time the patient had been treated with insulin 
for a total of three months.

The patient was kept in hospital for three further 
weeks and all the time he appeared normal as regards 
the blood and urinary sugars. The day before he was 
discharged there was a mean blood sugar level of 
130 mg./100 ml. and a delta blood sugar (\(= \Delta \text{B.S.}\)), 
\(i.e.,\) the difference between the maximum and the 
minimum values of the blood sugar curve, of 70 mg./ 
100 ml. At discharge the patient had increased his 
weight by 3.5 kg. and thus more than doubled his birth 
weight by the age of 4 months (Fig. 2).

Since his discharge in June, 1953, the patient has been 
examined several times. The blood sugar has been 
normal, there has been no glycosuria, and repeated 
glucose tolerance tests have proved normal too, most 
recently three years after discharge.

**Case 2.** A girl was born on April 1, 1955, at the 
Obstetric Department of the Centrallasarett in Våners-
borg. The parents were healthy, without any diabetic 
heredity. The mother was healthy during pregnancy. 
Confinement was normal and at term. The patient 
measured 47 cm. but weighed only 2,020 g., was strikingly 
thin, greyish-pale and sucked poorly. The weight dropped 
to 1,860 g. on the third day of life and then began to 
increase slowly.

On the eleventh day of life glycosuria and a urinary 
specific gravity as high as 1 029 was ascerained. A 
blood sugar test showed 400 mg./100 ml. and during the 
following three days the blood sugar rose to 440, 480 
and 560 mg./100 ml. An isolated test for urinary sugar 
at the age of 14 days showed 7.8%. No ketonuria was 
found. The patient was fed with breast milk in the 
usual quantities but fed poorly during the first two 
weeks.

At the age of 14 days insulin therapy was begun with 
6 i.u. IZS (Lente). In response to this the glycosuria 
disappeared, the specific gravity of the urine fell to about 
1.005, and the blood sugar began to fall. At the age of 
19 days the insulin dosage was increased to 8 units. 
Afterwards the patient did well except on one occasion 
when she showed an insulin reaction due to an alteration 
in the diet owing to a misunderstanding. The patient's 
weight increased quickly and at the age of 3 months she 
weighed 3,970 g.

Gradually the blood sugar tended to fall and simulta-
neously it became more stable, and the insulin dosage 
was reduced to 6 i.u. at the age of 37 days, to 4 units on 
the 57th day and finally was discontinued on the 200th 
day. The patient was kept at the hospital for another 
fortnight, and during this period the blood sugar was 
stable at a level of 90 to 160 mg./100 ml. A glucose 
tolerance test on the eighth day after discontinuing the 
insulin with 1 g. glucose per kg. body weight showed an 
increase of the blood sugar to 190 mg./100 ml. after 
75 minutes and a return to the level of 100 mg./100 ml. 
within three hours. A later test at the age of 1 year 
showed normal blood sugar levels and no urinary sugar. 
The child's length was 76 cm. and weight 10,160 g.; 
somatic and mental development were quite normal.
Discussion

It is open to discussion whether one is justified in calling these cases diabetic. Lawrence and McCance (1931) considered the diagnosis of diabetes mellitus justified if the patient showed diabetic symptoms and a glycosuria exceeding 2% or a clear recurrent hyperglycaemia above 200 mg./100 ml. Most of the above mentioned cases directly comply with these criteria, although Keidan's (1955) cases, for example, never showed pronounced hyperglycaemia. The fact that the diabetic condition was cured in these cases challenges a priori the diagnosis of diabetes mellitus. Joslin, Root, White and Marble (1952) have warned against exaggerated optimism, and claim a five-year period without recurrence as a criterion for recovery. In diabetes in children one might be justified in claiming curability even if the cases in question cannot claim a five-year period without recurrence. The three-year observation period of Case 1 of this report should be sufficient for regarding this case as recovered.

The absence of ketosis in these cases, at any rate, is remarkable, especially considering that the hyperglycaemia was often excessive. This may be explained partly by the type of diabetes in question here, and partly because a newborn infant seldom reacts with ketosis. As a general rule these cases of temporary diabetes have had but little need for insulin, and Keidan (1955) indeed asks whether insulin therapy really is necessary. In this connexion it may be said and emphasized that an attempt to discontinue it in our first case immediately resulted in aggravated conditions of the blood and urinary sugars. However, it is correct when Keidan (1955) points out that any greater insulin sensitivity is not always present initially. Because of these departures from the usual clinical picture of diabetes we would suggest the use of the term neonatal pseudo-diabetes mellitus.

An analysis of previously reported cases gives reason to suspect some of the patients to be postmature. Ramsey's (1926) case weighed 2,200 g. (about 5 lb.), Strandquist's (1932) and Arey's (1953) cases too weighed 2,200 g. each, and Keidan's (1955) patient 2,790 g. (6 lb. 2 oz.) though born at term. One of our cases weighed 2,780 g. and was clearly postmature, the other weighed 2,020 g. (about 4½ lb.) though born at term. It cannot be ignored that there has been postmaturity in a number of the cases described in the literature similar to our Case 1 where there were pronounced signs of postmaturity, i.e., a placental dysfunction or insufficiency.

Arey (1953) has postulated that in his case it was a question of a hyperadrenocorticism, and Keidan (1955) suggests a hypothalamic effect in his case. Much can be said in favour of the assumption that in these patients with neonatal pseudo-diabetes mellitus we are dealing with a carbohydrate disturbance of a steroid-diabetic character. In favour of this hypothesis is the occurrence of hyperglycaemia without ketosis, the relatively weak insulin sensitivity and curability. If in these cases, as at least in one of ours, a placental dysfunction led to postmaturity in the infant it is reasonable to presume a carbohydrate of a steroid-diabetic nature. It is known that the hormonal activity in the placenta even includes secretion of cortico-steroids of the '11-oxy' type. Priscilla White (1956) has thought that in these cases it is a stress or steroid type of condition.

Whether the alpha-cell dominance in the islets of the pancreas prevailing in an infant of this age—with the consequent hyperglucagonaemia—has any bearing on the occurrence of neonatal pseudo-diabetes mellitus cannot be decided. A discussion of this problem may even be of a theoretical nature only. The exact mechanism of these pseudo-diabetic cases thus so far is obscure and continual investigations are of the greatest value. If the urine is systematically examined in all infants during the neonatal period more cases would no doubt be recognized.

At present it cannot be said whether an actual diabetes mellitus will develop in these patients later on. However, it seems unlikely and the case reported by Ramsey (1926) at a follow-up examination 25 years later was normal. However, it may be advisable to make control tests in connexion with infections or other conditions that might act as a releasing factor for a latent diabetic disposition.

Summary

The authors report two cases of temporary diabetes during the neonatal period. Previously reported cases of the same nature are analysed and the term 'neonatal pseudo-diabetes mellitus' for this condition is suggested.

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