Occurrence of suppurative lymphadenitis after a change of BCG vaccine

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Abstract
BCG vaccines are the oldest vaccines in use today, but the protective effect of the vaccination is still controversial. The risk of contracting tuberculosis is low compared with the possible complications after this vaccination. In Austria the formerly used BCG vaccine was not available in the required amount and another vaccine was released by the drug authorities. This product, with a more virulent strain, was used between August and December 1990, and this increased the incidence of complications. Eighty four of 1950 vaccinated newborn babies developed severe suppurative lymphadenitis three to 28 weeks after the vaccination, and surgical treatment was found to be necessary. Isoniazid treatment did not prove to be successful when the lymph node exceeded a certain size. Culture was successful in 46% up to week 20; after 20 weeks no culture became positive. All cultured bacteria were isoniazid sensitive.

The question of general vaccination is raised and several points were considered before we came to the conclusion that except for high risk groups a general vaccination programme for neonates is not justified in Western countries.

BCG (Bacille Calmette-Guérin) vaccines are the oldest vaccines in use today. They were derived in 1906 by in vitro attenuation and, after 230 in vitro culture passages by Calmette and Guérin, were first used as oral vaccines in 1921. The subcutaneous application used at the present time was introduced by Wallgren in 1927.

After vaccination a positive tine test has been reported in 44% to 84% of cases and an even higher percentage (98%) reported for the Mantoux test two years after vaccination.1,2 The protective effect of the BCG vaccination has been estimated in many studies to be between 0 and 80%; in infants it seems to lie around 60%, based on known contacts of persons with confirmed tuberculosis.3,4 Recently the preventive impact of BCG vaccination has been shown by reports of high risks of tuberculosis in infants—in Egypt 76·7% and in Togo 64·5%—who had not been vaccinated.5,6 The protective effect of vaccination wanes significantly after the age of about 5 years.1,4–7

In 1952 the Ministry of Health in Austria, as in many other countries except for the USA and the Netherlands, both of which never used it on a national scale, recommended a general vaccination of newborn infants as a preventive measure against tuberculosis. In Germany the vaccination of the newborn was stopped due to complications in 1976. In Austria the risk of contact with tuberculous bacilli during infancy is low (the incidence of reported cases in the whole population is less than 0·05%) except in diseased families.8 Complications with BCG vaccines, such as suppurative lymphadenitis, osteitis, or even generalised tuberculosis, were reported in 0·01–3·8% even with the earlier strain. Nevertheless, in our hospital we had not performed any operation for suppurative lymphadenitis since 1974.9,10

Taking into account the low risk of acquiring tuberculosis (0·03%) and the incidence of severe complications from using the vaccine, the recommendation in Austria in 1989 changed to vaccination of high risk groups only: those in slum districts, refugee quarters, or endemic areas, children of diseased families, or of parents working in danger zones such as certain hospitals.11 Nevertheless, many paediatricians did not change their views and continued with general vaccination.

Based on the false assumption that paediatricians were following the health authorities, recommendation, BCG Berna (strain Kopenhagen) was not stored in a sufficient amount. The vaccine BCG-Pasteur Intradermal P, Charge R 5520, was released by the Austrian drug authorities and the usual demanding licencing procedures were dropped. However, this product contains a much more virulent strain of the tubercle bacillus. It was already known by the World Health Organisation (WHO) to produce a higher rate of complications,12 and for this reason the vaccine had been replaced. The Pasteur vaccine was used in some hospitals in the dosage of about 0·05 ml intradermally from the beginning of August to December 1990 until its use was prohibited by the Ministry of Health on the 28 November as a result of the alarming reports of a sudden increase in the occurrence of suppurative lymphadenitis.

The known fact led to the conclusion that the new vaccine raised the incidence of complications—a suspicion we initially expressed in a preliminary report.13

Subjects and methods
The department of paediatric surgery of the University Hospital Innsbruck is the only centre for paediatric surgery in the Tyrol, an area with about 500 000 inhabitants. In the critical period from August to December 1990, 3386 children were born and 1950 of these were vaccinated with the Pasteur vaccine. This is documented
by the authorities responsible for public health in the country. The dosage used was 0.05 ml, but in intracutaneous application the real amount is not exactly determined, because a small amount of the vaccine flows back due to the high intracutaneous pressure.

All children in this area were admitted to our hospital if surgical treatment was found necessary. The indication for surgical treatment was a fluctuating lymph node exceeding 1 cm in diameter, the inflammation infiltrating the surrounding skin, or spontaneous perforation or fistula. Under general anaesthesia an excision and excochleation of the necrotic tissue was performed. The material was sent to the Institute for Bacteriology where culture was developed on Löwenstein Jensen agar.14

Results
All 1950 newborn babies were mature except for one who was born in the 37th gestational week. The mean (SD) birth weight was 3500 (480) g. From November until 10 June, 84 of these children developed symptoms and were admitted for surgical treatment. The time between vaccination and surgery was surprisingly long. Suppurative lymphadenitis occurred within three to 28 weeks with a mean of 13.8 weeks (fig 1).

At the time they underwent surgery 34.5% of the infants had been on isoniazid treatment for one to 12 weeks already. Localised lymphadenitis was present in 44 boys and 40 girls: its distribution was axillary (59.5%), suprascapular (29.7%), nuchal (3.6%), cervical (3.6%), scapular (2.4%), and on the upper arm (1.2%). In 92.9% it was unilocular, and 7.1% had more than one enlarged lymph node at a different location. The size ranged from 1 to 4 cm in diameter. The area was typically reddened and in 4.7% spontaneous perforation occurred (figs 2–4). In 92.9% one operation, in 5.9% two, and in 1.2% three repeated operations were necessary due to further occurrence to lymphadenitis in another location.

In these infants the suppurative lymph nodes were totally removed surgically and showed caseous necrosis after incision. The excised necrotic tissue was sent to the Institute for Bacteriology for culture. On Löwenstein Jensen agar cultural detection of tubercle bacilli was successful in 46% up to week 20 after vaccination. After 20 weeks no culture became positive. All cultured bacteria were isoniazid sensitive.

All children were treated with isoniazid 10 mg/kg body weight plus 10 mg vitamin B-6 (Benadon, Roche) daily for six weeks.
Discussion

Infection from tuberculosis is rare in neonates and infants, but when it happens it is dangerous and means a prolonged morbidity despite all known therapeutic measures. In principle, a vaccination should solve these problems but a recommendation for general vaccination against tuberculosis needs some consideration.

Firstly, the incidence of tuberculosis has decreased dramatically. This started even before the tubercle bacillus was detected and was due to improvements in social standards and nutrition during the last 100 years. Neither chemotherapy nor vaccination alone have influenced this drop in cases of tuberculosis significantly. Therefore the number of infectious persons with open tuberculosis had declined. 8 11 13 16 Incidence lies now about 0·03% in Western countries.

Secondly, there are complications arising from the use of tubercular vaccines. The rate overall is estimated at about 0·01% to 3·6% comprising suppurative lymphadenitis (1/100), oesteitis (1/100 000), and even fatal generalised tuberculosis (1/1 000 000). 2 17-19 Reported complication rates depended on the dosage of vaccine used being 0·68% and 0·09% respectively for $2 \times 10^5$ and $0·5 \times 10^5$ viable units of the Pasteur vaccine. In striking contrast is our rate of severe complication with $1 \times 10^8$ viable units of this vaccine, making surgical treatment necessary in 4·5%. One report showed the non-suppurative lymphadenitis to occur in 18% of children vaccinated with a Pasteur vaccine. 20

Thirdly, the protection rate was supposed to be 80% for a duration of about 15 years, an opinion based on positive tine or Mantoux tests. However, there are reports on the waning effectiveness of vaccination after the age of 5 years. Even if in non-vaccinated children the rate of infection is higher than in vaccinated ones it is not acceptable to compare these two groups by randomised studies only without taking the social and nutritional status into account. 15 21 The most reliable report is a case-control study 22 showing a protection rate of 64%.

Fourthly, vaccination means a loss of the diagnostic value of intracutaneous tests (tine, Mantoux), which only show a positive reaction due to vaccination and not to 'wild type' infection.

Statistically, a life span of 100 years carries the same risk of acquiring tuberculosis as of suffering severe side effects due to vaccination. Even with vaccination there is no guarantee of total protection; the possibility of increased resistance has never been confirmed. Furthermore, no positive effect of vaccination in reducing the death rate can be shown; since 1970 no child (0–14 years) living in our area had died from tuberculosis. The tine test of Mantoux test is no proof of protection, it shows only that contact with bacilli has taken place.

After taking all these facts into consideration we must come to the conclusion that a general vaccination programme for neonates lacks justification. The WHO and also the Austrian health authorities met these deliberations with their recommendation to stop general vaccination in neonates except in 'high risk groups'.
The consequences of this occurrence, which raised hysterical reports and discussions in the press and led to a considerable financial burden for Austria, should result in preventive measures to keep the standards of medical services at a high level in the true sense of primum non nocere.


23 Texierieres L, Diesl MA, Chaud P, Saint-Cyr A, Salious P. Comparative trial of administration of half (0.05 mg) and quarter (0.025 mg) dose of intradermal Pasteur BCG on 291 infants from birth to 1 year in French Guyana. Vaccine 1991;9:521-4.