Nijmegen breakage syndrome

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Abstract
Nijmegen breakage syndrome (NBS), a rare autosomal recessive condition also known as ataxia telangiectasia (AT) variants V1 and V2, is characterised by microcephaly, typical facies, short stature, immunodeficiency, and chromosomal instability. We report the clinical, immunological, chromosomal, and cell biological findings in 42 patients who are included in the NBS Registry in Nijmegen. The immunological, chromosomal, and cell biological findings resemble those in AT, but the clinical findings are quite different. NBS appears to be a separate entity not allelic with AT.

Key words: Nijmegen breakage syndrome; ataxia telangiectasia.

Nijmegen breakage syndrome (NBS) is a rare autosomal recessive condition, characterised by microcephaly, typical face, short stature, chromosomal instability, and immunodeficiency accompanied by recurrent infections and predisposition to malignancies (MIM No. 251260). Since the recognition of the first two patients in 1981, we have included 42 patients in our NBS Registry in Nijmegen. The disease appears to be prevalent among the eastern and central European population, in particular among the Polish people. Although the immunological, cytogenetic, and cell biological findings in NBS resemble those in ataxia telangiectasia (AT), the clinical findings are quite different. NBS has been proposed as an allelic form of AT and designated as an AT variant with two complementation groups, V1 and V2.

Recently the gene for AT on chromosome 11q22-23 has been identified (ATM=AT mutated) and found to be mutated in AT patients from all four complementation groups. At present there is little information on the possible chromosomal location of the NBS gene(s). The AT locus, however, has been excluded from harbouring the NBS gene.11,13-15

Clinical description
GROWTH AND DEVELOPMENT
The growth retardation in NBS patients can be of pre- or postnatal onset. In 70% of patients, birth weight and birth length conform to gestational age. The head circumference (OFC) at birth varies between 26-5 cm and 36 cm; 75% of patients are born with an OFC below the 3rd centile. This implies that not all patients are microcephalic at birth. However, they all develop progressive and severe microcephaly during the first months of life. Height falls below the 10th centile in all patients. From the first stages of development onwards growth retardation is observed. After a few years a slight improvement is seen and height is on average almost on the 3rd centile. The growth retardation is proportionate and weight corresponds to length. Bone age can be somewhat retarded. Little is known about the sexual maturation of patients with NBS. The only endocrinological abnormality described as yet is ovarian dysgenesis in two patients (K Chrzanowska, personal communication).

Developmental milestones are mostly not delayed during the first year of life. Mental development is normal in 35% of NBS patients; 45% have borderline retardation, while 20% are moderately retarded. None of the patients has signs of cerebellar ataxia, apraxic eye movements, or other neurological abnormalities, except for two girls described by Curry et al, who had clinical symptoms of both NBS and AT. Their condition was considered to be an AT variant and referred to as “ATFresno”, representing a distinct entity. Complementation studies performed in 1988 assigned these twin sisters to NBS complementation group V1.7

CRANIOFACIAL FINDINGS
The craniofacial characteristics become more obvious with age. Although the progressive and severe microcephaly in all patients. The hair becomes sparse in half of the NBS patients. The face is characterised by a receding forehead, prominent midface with a long nose and long philtrum, receding mandible, upward slanting palpebral fissures, large ears with dysplastic helices, and freckles on the cheeks and nose (figs 1 and 2). Subtle scleral telangiectasia can be present; this has been noted in 10 out of 25 patients.37

SKIN
Pigmentation abnormalities are present in most patients. Freckles are mostly seen on the face.
Two to five café au lait-like spots are found in half of the patients and most also have some vitiligo spots. Cutaneous telangiectasia have been noted in only two patients. Sun sensitivity of the eyelids can be present.

CONGENITAL MALFORMATIONS
Congenital malformations reported once each are: preaxial polydactyly, occipital cyst, hydrocephalus, choanal atresia, cleft lip and palate, hypospadias, and a single ectopic kidney. In three patients anal atresia was present at birth, and in another three patients hydrocephalus has been diagnosed.

INFECTIONS
Respiratory tract infections occur in almost all patients with NBS. Pneumonia, bronchitis, otitis media, sinusitis, and mastoiditis are frequently seen. Complications such as bronchiectasis, respiratory insufficiency, and even death from respiratory failure can occur. In three patients no respiratory tract infections were reported, but two of them were only 8 and 18 months old at the time of reporting. The other patient is a 4 year old girl who had no disturbance of her cellular immunity. Urinary tract infections repeatedly occur in 15% of patients with NBS. Gastrointestinal infections with diarrhoea have also been mentioned in 15% of patients.

PREDISPOSITION TO MALIGNANCIES
Both immunodeficiency and chromosome instability probably predispose to malignancies. In NBS patients there is a tendency to develop lymphomas at a young age. So far, 12 patients varying in age from 1 to 22 years have developed a lymphoma. One patient developed a glioma at the age of 12 years, one patient a medulloblastoma at 15 years, and one patient a rhabdomyosarcoma at 4 years. Two patients who developed a lymphoma are still alive, the others died within a few months to years after the malignancy had been diagnosed. When treating malignancies in these patients, cytostatics should be the first choice. However, radiomimetics (for example, bleomycin) should be avoided, and the chemotherapy doses should be reduced. Radiation therapy should also be avoided, since irradiation can induce malignancies in NBS patients. If necessary, as can be the case in solid tumours, an estimation of the individual dose is recommended.

NECROPSY FINDINGS
Of the 42 patients included in the NBS registry 15 have died, four patients from bronchopneumonia, 10 from lymphoid malignancy, and one patient who developed a medulloblastoma. An extensive necropsy was performed on only one patient. In this patient important findings were the extremely reduced brain weight and the absence of diffuse cortical cerebellar degeneration, the presence of which is characteristic of AT. The thymus showed
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Immunological disturbances
The immune deficiency in NBS patients is relatively severe and concerns the humoral and cellular immune system. The humoral immunodeficiency is characterised by disturbed serum immunoglobulins; agammaglobulinemia, IgA deficiency, and IgG2 and IgG4 deficiency can be found. In one patient immunological attrition was noticed. As well as IgA deficiency that he had had for years, he developed agammaglobulinemia at 24 years of age, after he had been treated with cytostatics for a lymphoma. The defective cellular immunity is characterised by mild to moderate lymphopenia with decreased CD3+ and CD4+ (helper) cells and a decreased CD4+:CD8+ (suppressor) cell ratio. The in vitro response of lymphocytes to mitogens is impaired. The patient described by Barbi et al was the only one reported to have normal T cell immunity and had no recurrent infections at the age of 4 years.

Cytogenetic features
NBS patients basically have a normal karyotype. Cultured T lymphocytes often show an extremely low mitotic index, in which case cytogenetic analysis can be very difficult. In all patients structural chromosome aberrations are present in 10 to 35% of metaphases of cultured T cells. Most of the rearrangements preferentially occur in chromosomes 7 or 14 or both at bands 7p13, 7q35, 14q11, and 14q32, the same breakpoints as found in AT. These breakpoints coincide with the location of genes having immune function. Inv(7)(p13q35) is the most frequently detected aberration in NBS. Other rearrangements are t(7;14) (p13;q11), t(7;14) (q35;q11), t(7;7) (p13;q35), and t(14;14) (q11;32).

Inv(14)(q11q32) had not been found in NBS patients until recently when it was detected in a lymphocyte culture of a 24 year old patient. In fibroblasts, chromatic type instability has been observed. In one case chromosomal rearrangements were found in 33% of fibroblasts, but without involvement of chromosomes 7 or 14. No chromosomal instability has been observed in bone marrow cells.

Alpha fetoprotein (AFP) levels
Serum AFP levels are normal in NBS patients, in contrast to AT patients, in whom an increased serum AFP level is one of the hallmarks. The only patients with abnormally increased serum AFP levels are the twins with ATFresno, assigned to NBS complementation group V1. In two patients, serum AFP levels were raised at the age of 5 months, but decreased to normal by 15 months and 2 years respectively.

Radioresistant DNA synthesis (RDS) and radiation hypersensitivity
Increased sensitivity of both lymphocytes and fibroblasts from NBS patients to ionising radiation or to radiomimetics, and radioresistance of DNA replication, have been established. So far, all patients tested have shown an abnormally increased rate of cell death and chromosome damage after x or gamma irradiation. The inhibition of DNA synthesis after x or gamma irradiation is two to three times less pronounced than in normal cells. The results are indistinguishable from AT.

Genetics and molecular studies
NBS is an autosomal recessive condition. It has been diagnosed in at least 42 patients, 23 males and 19 females from 29 families. The NBS Registry contains 12 sibships among the 42 patients included.

NBS seems to be more prevalent among the Polish people. Twenty-one of the 42 patients have been diagnosed by one of the authors (KH C) in Warsaw. The aetiology of NBS is still completely unknown. Because the immunological, cytogenetic, and cell biological findings in NBS are almost the same as in AT, the elucidation of the AT gene would be an important step forwards for unravelling the genetic basis of NBS. Although NBS proved not to be allelic to AT, the gene responsible for NBS may be similar to the ATM (=AT mutated) gene and have comparable functions. The ATM gene on chromosome 11q22-23 was found to be mutated in AT patients from all complementation groups. A partial ATM complementary DNA clone encoded a putative protein with sequence similarities suggesting involvement of the ATM gene product in signal transduction, cellular responses to DNA damage, and cell cycle control functions that are defective in AT.

The observation of non-linkage of NBS with the ATM gene was corroborated by the study of Komatsu, who found non-complementation of NBS fibroblasts with chromosome 11.

Hitherto, NBS patients have been divided into two different complementation groups, V1 and V2, whereas AT patients have been represented in four other separate groups: A, C, D, and E. The characterisation of the ATM gene and convergence of all complementation groups into one gene requires reconsideration of the complementation groups in NBS. NBS is probably also a homogeneous disorder. Knowledge about the ATM gene products and their function could provide important clues for the aetiology of NBS.

Prenatal diagnosis
As the aetiology of NBS is still unknown, possible methods of prenatal diagnosis must depend on the demonstration of the biochemical findings of the disease. The assessment of spontaneous or radiation induced chromosome aberrations in prenatal tissue has been reported in only a few AT cases. The disadvantage of
this method is that a large number of mitoses in chorionic villus (CV) or amniotic fluid (AF) cells have to be analysed. Radiodistant DNA synthesis (RDS) assay on cultured CV or AF cells has proven to be a reliable method for first trimester prenatal diagnosis of NBS and of AT. The prerequisite for reliable prenatal diagnosis with this method is the demonstration of increased RDS in fibroblasts of the index patient.25

**Differential diagnosis**

The well known chromosome instability disorders are Fanconi anaemia, Bloom's syndrome, and ataxia telangiectasia. Fanconi anaemia is characterised by short stature, pancyclopenia, radial hypoplasia, and brownish hyperpigmentation. Important hallmarks of Bloom's syndrome are severe growth retardation of prenatal onset, sunlight sensitivity leading to telangiectatic erythema, and specific chromosome instability with a high frequency of sister chromatid exchanges. Common findings of both AT and NBS are immunodeficiency, chromosome instability with multiple chromosome 7 or 14 rearrangements, chromosomal hypersensitivity to x-rays, and radiodistance of DNA replication. The classical clinical findings in AT are progressive cerebellar ataxia, ocultocutaneous telangiectasia, and raised serum AFP levels, which are absent in NBS.

**Conclusion**

Because patients with NBS have an increased risk of developing malignancies, it is important to diagnose this disease at an early age. In NBS patients x irradiation should be avoided as much as possible, for therapeutic but also for diagnostic reasons.

In all patients with unexplained early progressive microcephaly and growth retardation, chromosome instability analysis should be performed, in order to establish or exclude a chromosome instability disorder.

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