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Spontaneous pneumothorax as a clinical manifestation of neurofibromatosis type 1

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SUMMARY
Neurofibromatosis type 1 (NF1) is a genetic disorder affecting the skin, nervous system, eyes and bones. Pulmonary involvement is unknown to many physicians. Yet, patients may be affected by lung bullae and cysts, which represent an increased risk for secondary spontaneous pneumothorax (SSP). We present a 56-year-old patient with a pathogenic variant of the NF1 gene, who suffered from NF1 with lung manifestations and recurrent SSP. It is essential to identify the patients having an increased risk of developing SSP as preventive surgery seems to decrease the risk of new events. Pneumothorax can be a clinical manifestation of NF1 but is not yet widely acknowledged as such.

BACKGROUND
Neurofibromatosis (NF) is a condition which affects the skin, eyes, lungs, bones and nervous system. NF is classified in three heterogeneous groups: NF1, NF2 and schwannomatosis. NF1, also known as von Recklinghausen’s disease, is the most common form with a prevalence of approximately 1 per 3000.1 NF1 is an autosomal dominant disease caused by a mutation of the NF1 gene. The mutation is, however, not a diagnostic criterion. The diagnosis is based on the presence of two or more of the clinical features presented in box 1.

Thoracic manifestations are most commonly cutaneous and subcutaneous neurofibromas on the chest wall and kyphoscoliosis.3 Skeletal manifestations include deformities of the ribs, posterior vertebral scalloping and enlargement of the intervertebral foramen. Many physicians are not aware that up to 10%–20% of the patients have involvement of the lungs.4 The pulmonary manifestations include apical lung bullae and cysts, thinned bullae, interstitial parenchymal lesions and NF-related interstitial lung disease with bilateral, symmetrical and basal involvement of the lungs. Intrathoracic neurogenic tumours, meningoceles and plexiform neurofibromas of ribs, chest wall, lungs and mediastinum have also been described.

CASE PRESENTATION
A 56-year-old man diagnosed with NF1 was referred to our clinic with a suspicion of Birt-Hogg-Dubé syndrome (BHD) due to a history of recurrent spontaneous pneumothorax. He was diagnosed with NF1 28 years earlier. The physical examination of the patient showed multiple café-au-lait macules, freckling of axillae and groins, multiple neurofibromas and plexiform neurofibromas on his face, trunk and extremities, and fibromas of the tongue. The patient was also known to have an optic pathway glioma in his right eye. He had a mother and a sister with known NF1. He thereby met five of the diagnostic criteria of NF1 including first-degree relatives with NF1.

He was a former smoker with a smoking history of 10 pack years. During the years, he had seven neurofibromas and one malignant schwannoma removed. Furthermore, he had a variety of other health issues including Barrett’s oesophagus, irritable bowel syndrome, bilateral sensorineural hearing loss and conductive hearing loss on his right ear.

He had a remarkable history of at least four incidents of pneumothorax. Both lungs had been involved, and each pneumothorax was treated by tube thoracostomy. The most recent incident was in 2006. The patient attended his general practitioner as he experienced shortness of breath and pain located to the left side of the chest. An X-ray showed an apical pneumothorax of the left lung. The pneumothorax was treated by tube thoracostomy for two days, until the lung was fully inflated. He was referred to the Department of Thoracic Surgery as several bilateral lung bullae and apical cysts were demonstrated on a high-resolution computed tomography (HRCT) (figure 1). However, the cysts were not removed surgically due to the diffuse localisation. Furthermore, it had been 10 years since the previous incidence of secondary spontaneous pneumothorax (SSP). It was argued in the medical record that the operation would not benefit the patient, and the risk of doing harm was too great. The same reason was given, when a chest scan in 2017 revealed several cysts, bilateral paraseptal and centrilobular emphysema, pleural thickening, and fibrosis causing interstitial lung disease.

The patient was genetically tested for NF1 and BHD, and was found to have a pathogenic variant only in the gene encoding NF1 (c.6709C>T; p.Arg2237).

Box 1 Clinical features of neurofibromatosis type 1 (NF1)

► Six or more café-au-lait macules
► Two or more neurofibromas, or one or more plexiform neurofibroma
► Skinfold freckling of axilla or groin
► Optic pathway glioma
► Two or more Lisch nodules
► Characteristic bony dysplasia (sphenoid wing dysplasia, thinning of the long bone cortex)
► First-degree relative with NF1

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**Case report**

Figure 1  HRCT from 2007. The scan is showing apical lung cysts and paraseptal, bullous and centrilobular emphysema. Areas without emphysema or lung cysts are affected by ground glass opacities.

**OUTCOME AND FOLLOW-UP**

The Centre of Rare Diseases is following the course of the patient’s disease by regular follow-ups. The patient has not had any worsening in his pulmonary aspects, although he does suffer from shortness of breath.

The patient is still seen at the Department of Dermatology for follow-up. The treatment consists of CO₂ laser to reduce the neurofibromas and surgery to remove deeper neurofibromas causing him pain and discomfort.

**DISCUSSION**

In this case, BHD was considered by the clinical geneticist due to the recurrent episodes of SSP. In about 5%–10% of patients with spontaneous pneumothorax, the underlying disorder is BHD, which is characterised by lung cysts, renal tumours and fibrofolliculomas. BH D is caused by a mutation of the FLCN gene of which our patient was tested negative. Other genetic diseases with pulmonary involvement and possible pneumothorax include tuberous sclerosis, Ehlers-Danlos syndrome, Marfan syndrome, Langerhans cell histiocytosis and NF1 of which our patient was affected. Neither the pulmonologists nor the thoracic surgeons associated the pneumothorax with the patient’s NF1.

Differential diagnoses of NF1 include NF2, Legius syndrome, familial café-au-lait macules, Noonan syndrome and constitutional mismatch repair-deficiency syndrome. NF1 is genetically and clinically distinct from the above mentioned, but especially the neurofibromas can be mistaken for other types of tumours. Genetic testing can therefore help to differentiate NF1 from other diseases. The diagnosis of our patient was confirmed by a disease-causing mutation of the NF1 gene. Due to his two first-degree relatives with NF1, the mutation seems likely to be inherited, although none of the family members have been genetically tested. The patient also had a son, who had spontaneous pneumothorax at the age of 17, but lacked other clinical features of NF1.

To our knowledge, this is the first reported case with a pathogenic variant of the NF1 gene and SSP. In the literature, we found one case reporting spontaneous pneumothorax as the initial presentation of NF1. To investigate the association between NF1 and pneumothorax, a PubMed search was carried out. This revealed 10 published cases of pneumothorax in NF1 (Table 1).

All but one of the published cases had pulmonary manifestations of NF1, besides pneumothorax, and two of the cases were known non-smokers. Smoking might increase the risk of interstitial lung disease, but the incidence of SSP is independent of smoking habits. The literature seems to agree that cystic, bullous or emphysematous lung disease are distinct clinical manifestations of NF1. None of the cases reported NF-associated interstitial lung disease characterised by pulmonary fibrosis or symmetrical, basal affection of the lungs. Thus, the occurrence of SSP is likely a clinical feature directly associated with the cystic and bullous lung manifestations of NF1 itself, and not a result of the NF-associated interstitial lung disease. Thereby, pneumothorax may be directly caused by NF1.

Correlation between genotype and phenotype in NF1 is not yet well known due to gene size, phenotypic complexity,

| Table 1  Reported cases of pneumothorax in patients with neurofibromatosis type 1 (NF1) |
|---------------------------------------------------------------|
| **Clinical features** | Ayed et al⁹⁹ | Engdahl et al⁹⁸ | Gangadi et al⁶⁶ | Nardecchia et al¹¹ | Nguyen et al¹⁰ | Satar et al¹¹ |
| Cafe-au-lait macules and spontaneous pneumothorax | Cafe-au-lait macules, soft tissue masses, and spontaneous pneumothorax | Neurofibromas and spontaneous pneumothorax | Cafe-au-lait macules and spontaneous pneumothorax | Cafe-au-lait macules, inguinal freckling, neurofibromas, phaeochromocytoma, and spontaneous pneumothorax | Axillary freckling, cafe-au-lait macules, bilateral Lisch nodules, neurofibromas, first-degree relative with NF1, mixed-type hearing loss, and spontaneous pneumothorax |
| **Radiology** | Apical, paraseptal and parenchymal lung bullae, paraseptal emphysema, apical pleural thickness, and subpleural nodules | Spinal and paraspinal tumours | Centrilobular emphysema, subpleural thin-walled lung cysts, bilateral basal reticular opacities, basal linear atelectasis, and subcutaneous nodules on the chest wall | Bilateral apical lung bullae and emphysema | Asymmetric pulmonary oedema, paraseptal and centrilobular emphysema, ground glass opacities, and pneumatoceles | Air cysts in upper lobes |
| **Smoking status** | Non-smoker | N/A | Non-smoker | N/A | Smoker (50 pack years) | Smoker (33 pack years) | Smoker |

The PubMed search for NF1 and pneumothorax gave four additional hits. These were not included due to other languages than English, German or French. N/A, not applicable.
heterogeneity of genotype and phenotype, and phenotypic progression with age. In the literature, only few correlations have been identified. It is worth noticing that Satar et al reported a case of first-degree family history of NF1 and clinical features similar to our patient. Both had skinfold freckling, café-au-lait macules, multiple neurofibromas and mixed-type hearing loss as well as recurrent pneumothorax. This is the only other case besides ours to report recurrent pneumothorax and first-degree relatives with NF1. As none of the reported cases in table 1 were genetically tested, it is uncertain whether the pneumothorax of the patients was linked to genotype. Thus, a possible genotype–phenotype correlation between SSP and NF1 cannot be directly addressed.

Spontaneous pneumothorax is painful for the patients and can be lethal in 4.6% of the cases. It is therefore important that the clinician is attentive to the pulmonary manifestations of NF1 characterised by lung cysts and bullae. It is essential to identify the patients having an increased risk of developing SSP as preventive surgery seem to decrease the risk of new events.

Our case represented a male who met five of the diagnostic criteria of NF1 with pulmonary manifestations including several lung cysts and bullae and at least four incidents of SSP. This should lead to consideration of NF1 as the aetiology of pneumothorax incidents. The diagnosis of NF1 was confirmed by genetic testing showing a pathogenic variant of the NF1 gene. To address a genotype–phenotype correlation between disease causing variants of the NF1 gene and pneumothorax, genetic testing and screening for pulmonary manifestations in patients with NF1 are recommended for future clinical studies. Clinicians need to be aware that pneumothorax can be a manifestation of NF1.

**Learning points**

- Spontaneous pneumothorax is a clinical presentation of neurofibromatosis type 1 (NF1) caused by lung cysts and bullae.
- NF1 should be considered in patients presenting with spontaneous pneumothorax.
- It is of great importance that the clinicians are attentive about pulmonary manifestations in patients diagnosed with NF1.
- Early diagnosis of pulmonary manifestations in patients with NF1 is crucial as surgical removal of lung cysts and bullae seem to prevent spontaneous pneumothorax.
- An examination of genotype–phenotype correlation between NF1 and pneumothorax could be of interest to identify patients at risk.

**Contributors** TL wrote the first draft and edited the final version of the manuscript. AB commented, cowrote and edited the final version of the manuscript. HM edited the patient’s chest scans and revised the manuscript. MJZ-L-T provided the patient’s chest scans and revised the manuscript. AB was involved in the care of the patient.

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