For debate

Pulmonary sequestration and related congenital bronchopulmonary-vascular malformations: nomenclature and classification based on anatomical and embryological considerations

BS CLEMENTS, JO WARNER

From the Brompton Hospital, London

No satisfactory classification for congenital anomalies of the bronchopulmonary airway and related vasculature has yet been described. An attempt was made to overcome this problem by expanding the concept of "sequestration" (originally intended as a single entity) to include a range of bronchovascular anomalies. This has failed to gain universal acceptance and led us to develop an alternative nomenclature and classification based on simple anatomical description. This paper discusses the background to the new approach while a companion paper applies it to our own cases reviewed during five years.

History and background

Defining a congenital anatomical lesion of the lung affecting a single structure (for example, bronchus) is relatively simple (for example, bronchial stenosis, bronchogenic cyst). When this lesion includes abnormalities of other major components of lung tissue—namely, arterial supply, venous drainage, and lung parenchyma—differentiation becomes more complex. In 1946 Pryce coined the term sequestration to describe a "disconnected" ("dislocated," "ectopic") bronchopulmonary mass or cyst with an anomalous systemic artery supply.1 Four of his original seven cases differed considerably from his own definition, which led him to describe three variants of sequestration—namely: abnormal artery to normally connected lung; abnormal artery to both sequestered mass and adjacent normal lung; and abnormal artery confined to the sequestered mass.2 Later the terms intralobar and extralobar were added but they provide no definitive information on the morphology within these lesions. Despite this, many authorities have adhered to Pryce's terminology and consequently encountered problems in defining the variants of sequestration.3–6 Many synonyms for the various atypical forms have been proposed, contributing further to the confusion concerning nomenclature.7 Realisation that these entities may represent variants of a single primary complex of bronchovascular anomalies8–10 led to the concept of a "sequestration spectrum," a term introduced by Sade in 1974.10 This extended term included various combinations of abnormal bronchial connection, arterial supply and venous drainage, so bringing in the scimitar syndrome and its variations.11 In 1984 Thilenius again reviewed the sequestration spectrum concept.12 He emphasised the wide range of anomalies found by quoting examples of every combination of abnormal bronchial connection, arterial supply, and venous drainage appearing in published reports. He agreed with Sade, and others,13 that sequestered areas of lung with normal vasculature should be included in the spectrum. Nevertheless, reports of "atypical" sequestrations continued to inundate published reports,14–15 indicating that this concept had failed to gain general acceptance. To complicate matters further, associated abnormalities of venous drainage and lung parenchyma were rather neglected in most earlier reports. Recently the importance of defining the venous drainage in these lesions has been emphasised,12–16 and our experience further emphasises this aspect (see p 409). Associated abnormalities of lung parenchyma, both within the lesions and in the surrounding lung, have not been widely appreciated, although their existence was noted in Pryce's
original articles. We also expand on this feature in our clinical review.

From these descriptions it is apparent that there are essentially four components of these lesions—namely, the tracheobronchial airway, arterial supply, venous drainage, and lung parenchyma. We should therefore be able to construct a definition and classification based on these components that will cover all possible anatomical combinations. In this paper we propose a new nomenclature and classification with supporting evidence from histological and embryological observation. The companion paper (p409) illustrates the clinical advantages of this simple anatomical approach.

NOMENCLATURE AND CLASSIFICATION

There are several major objections to the use of “sequestration” as a collective term for all bronchopulmonary-vascular anomalies. These include:

1 The Concise Oxford Dictionary defines sequestration (from the Latin sequestrare, meaning among other things to remove or separate) as to seclude, isolate, set apart. This term is therefore incorrectly applied to the lesions previously described in published reports as, strictly speaking, only “extralobar sequestrations” are truly “set apart” from the normal lung.

2 The limits of the sequestration spectrum have never been defined; thus the range of anomalies included varies considerably according to the author.

3 Most people continue with the impression that “sequestration” is a “disconnected bronchopulmonary mass or cyst with a systemic arterial supply,” albeit fully accepting how nebulous this may be. Thus there is a natural reluctance to include lesions with normal vasculature in this category.

On the other hand, Stedman’s Medical Dictionary (1982) defines malinosculation (Latin: mal—in abnormal; in—in; osculum—mouth) as “the establishment of (abnormal) communications by means of small openings or anastomoses, applied especially to the establishment of such communications between already existing blood vessels or other tubular structures that come into contact.” This definition describes the pathogenesis of these congenital lung lesions exactly. If we are going to classify them accordingly, we must use this appropriate term.

Thus pulmonary malinosculation simply describes a congenitally abnormal connection or opening of one or more components of the bronchopulmonary vascular complex. The various combinations of these components give rise to a wide spectrum of abnormalities. The lesions thus formed can then be classified simply according to their abnormal anatomical components. The classification is constructed in three steps. The first step describes the abnormality of bronchopulmonary airway or arterial blood supply or both. The possible combinations of these two com
Pulmonary sequestration and related congenital bronchopulmonary vascular malformations

ponents form a spectrum of lesions, as illustrated in figures 1-3. At one end of the spectrum is absence or abnormality of bronchopulmonary airway with normal pulmonary artery blood supply (fig 1); this is bronchial pulmonary malinosculation or bronchopulmonary malinosculation. At the other end of the spectrum (fig 2) is an anomalous arterial supply to an area of lung with a normal bronchopulmonary airway; this is arterial pulmonary malinosculation. In between we have all the combinations and gradations of abnormal bronchial and arterial connection. These represent bronchoarterial pulmonary malinosculations (fig 3).

The second step in the classification defines the associated anomalies of venous drainage. The classification is then completed by describing any associated abnormalities of lung parenchyma. Each of these steps will now be reviewed in more detail.

TRACHEOBRONCHOPULMONARY MALINOSCULATION

Disruption of normal airway communication can occur at any level from the trachea down to the alveolus. Disruption may be incomplete (for example, stenosis) or complete (for example, atresia). The lesion may be localised or extend distally to affect a variable segment of the bronchopulmonary tree. These lesions are conveniently divided into those affecting proximal major airways (tracheobronchopulmonary malinosculation), such as tracheal stenosis and tracheal cysts, and those affecting the distal airways down to respiratory bronchioles and alveoli (bronchopulmonary malinosculation). This latter group includes lesions such as bronchial stenosis and bronchogenic cysts, and parenchymal lesions with normal pulmonary artery blood supply, such as congenital lung cysts, cystic adenomatoid malformation, and lobar emphysema. Each of these entities may have associated anomalies of venous drainage and lung parenchyma.4-12

ARTERIAL PULMONARY MALINOSCULATION

Arterial pulmonary malinosculation refers to an aberrant systemic arterial supply to an area of otherwise normal lung.1-7 Most of these lesions, however, also affect the tracheobronchopulmonary airway to some extent, which will mean that they are classified in the following category.

BRONCHOARTERIAL PULMONARY MALINOSCULATION

Here there is an abnormality of both the bronchopulmonary airway and the arterial blood supply to an area of lung. The abnormality of bronchopulmonary communication may range from absent connection with a blind ending bronchus (agenesis) or atretic membrane to apparently normal patency with a gradation of abnormality in between.4-9,10,12 Abnormal branching patterns, reduced numbers of bronchial branches, or generalised hypoplasia may also occur.12,18 The aberrant arterial blood supply can arise as one or more trunks, usually from the abdominal or thoracic aorta.3-11 An abnormal area of lung sharing a mixed pulmonary and systemic arterial blood supply has been described (see paper on p409). In addition, part of the surrounding normal lung may be supplied by the anomalous artery.1

Entities found in this group include sequestrations, lung cysts with aberrant systemic arterial supply, and congenital cystic bronchiectasis. Again, each of these lesions may have associated anomalies of venous drainage or lung parenchyma, or both (for example, scimitar syndrome).18

Abnormalities of venous drainage

Anomalous venous drainage can occur with any congenital lung malformation whether there is an aberrant systemic arterial supply or not (fig 1-3).12 The anomalous veins may be single or multiple, and drain directly to the right atrium, the major draining veins in the thorax or abdomen, or to both the right and the left atrium.16 Obstruction to venous return is said to occur in some cases.18 In one group of patients, particularly where the lesion is found in the right lung, the anomalous vein or veins drain the major part or the whole of that lung, while the aberrant arterial supply is limited to the right lower zone. This mismatched anomalous venous drainage is a characteristic feature of the scimitar syndrome (fig 4).12,18

Abnormalities of lung parenchyma

Here the site of the lesion is described, together with the changes occurring within the lesion (for example, cystic, adenomatous, emphysematous, or containing

"Mismatched" anomalous venous drainage

Fig 4 Bronchoarterial pulmonary malinosculation with mismatched anomalous venous drainage—a well recognised variant of the scimitar syndrome.
foregut inclusions) and associated abnormalities of surrounding lung parenchyma (for example, hypoplasia and abnormal lobation). The site of the lesion may be intrapulmonary (within the boundary formed by the pleural layer surrounding the lung) or extrapulmonary (ectopic lung tissue lying outside the boundary formed by the pleural layer surrounding the rest of the lung).

The classification of pulmonary malinosculation is summarised in table 1. The distinct clinical entities within the classification should all retain their common nomenclature. The purpose of this classification is not to replace their accepted terminology but to aid categorisation of these lesions, particularly the more complex varieties. In turn, it is hoped that this will improve the understanding and management of these complicated malformations. A good example of a complex lesion affecting all four major components of lung tissue is a well recognised variant of the scimitar syndrome usually presenting in infancy.16 18–20 This lesion is situated in the lower half of the right lung with abnormal lobation and bronchial branching, anomalous systemic arterial supply and venous drainage, and associated lung hypoplasia. A characteristic feature is the “mismatch” between the aberrant systemic arterial supply to the right lower zone and the anomalous venous drainage of the whole or major portion of the right lung (fig 4).18–20 According to our terminology, this malformation would be classified as bronchopulmonary pulmonary malinosculation with mismatched anomalous venous drainage and ipsilateral lung hypoplasia. In a clinical context, each component would then need to be further defined—for example, in terms of the degree of disruption in airway communication and the site of anomalous venous return. The information, with the clinical history and haemodynamic data, is essential for appropriate patient management.

**Discussion**

Application of a unifying concept to congenital lung malformations is certainly not new. In 1974 Buntain suggested that lobar emphysema, cystic adenomatoid malformation, pulmonary sequestration, and bronchogenic cysts in children were a clinical group.2 Heitzman went even further, defining the sequestered lung with normal pulmonary artery supply as “a small area of normal lung with systemic artery supply, much as we have described.10 Heitzman divided five of these malformations previously classified elsewhere—such as lobar emphysema, congenital cystic bronchiectasis, and cystic adenomatoid malformation.13 He warned that although the terminology suggests that these are discrete entities, their morphological characteristics overlap to a considerable degree. Demos removed from a five month old infant a lobe whose tissue, both grossly and microscopically, showed characteristics of “intra- and extralobular sequestration, bronchial atresia and cystic adenomatoid malformation.”22 As early as 1969 Culiner, when comparing cases of “congenital bronchial cystic disease, sequestration and adenomatoid cystic bronchiectasis,” suggested that these three entities were variants of a single primary complex of bronchovascular anomalies.8 It would certainly assist our understanding of their pathogenesis.
Pulmonary sequestration and related congenital bronchopulmonary vascular malformations

if they could be regarded in this way.

Fundamental to the subject of all malformations is the embryology of the part concerned. Unfortunately, no generally accepted explanation for the development of these lung anomalies has yet been suggested, despite 40 propositions in Biancalana's paper and the six most widely publicised theories. We propose a rational sequence of events in lung development, which would account for all lung anomalies that include malinosculation of one or more components of the bronchopulmonary-vascular complex, starting with a few widely accepted observations concerning embryogenesis, namely:

1. Bronchial branching is completed 16 weeks after conception.
2. Pulmonary artery development is similar to the pattern of bronchial branching but follows slightly later.
3. Alveolar development follows transformation of the terminal bronchioles into respiratory bronchioles and, later, saccules after about the 24th week from conception.
4. In addition to the recognised stages of lung development, there are differing rates of growth occurring in various tissues at different times.
5. In the early stages the tips of the dividing bronchi begin to develop more extensively from the 12th week onwards.

Fig 5 The "wheel" theory of abnormal lung development. After a theoretical "insult" to the tip of a dividing bronchus (shown in the centre of the diagram), each satellite sketch represents a possible pathological consequence at the next stage of development. This forms the basis of the eventual lesion.
chial buds are supplied by a systemic capillary plexus derived from the primitive aorta. This plexus regresses as the growing lung advances, the developing pulmonary artery taking over. 25 29 31

The pathological options that follow an insult to the tip of a developing bronchus can be graphically represented by the spokes of a wheel (fig 5). The "insult" is undetermined and could take the form of localised trauma, ischaemia, infection, adhesions, or any non-specific injury. It is not only the nature of the insult, however, but also—and probably more importantly—the timing and severity that determine the morphology of the eventual lesion. 30

The first possibility (1 in fig 5) is that development of the bronchus and its pulmonary artery supply is totally arrested, leading to agenesis of that lung, lobe, lobule, or segment. The timing will determine how distal the lesion is.

A second possibility (2 in fig 5) is that the insult may be minor, resulting in a localised abnormality followed by normal development of the distal bronchial tree, lung parenchyma, and pulmonary artery. This will result in a local lesion, such as bronchial stenosis or a bronchogenic cyst.

A third possibility (3 in fig 5) is that there is no interference with continued development of the bronchial tree, but pulmonary artery growth is arrested. Devoid of pulmonary blood supply, continued growth of the developing bronchial branch could be supported only by vessels retained from its normally regressing systemic capillary network. Growing along with the developing bronchial tree, these systemic vessels would become substantial channels, single or multiple, and establish a systemic arterial supply to the area of lung concerned. The sites of origin of these vessels move distally with the caudal "shift" of the growing primitive aorta from which they arise, ending up in the lower thorax or upper abdomen. Such an anomalous area of "normal" lung with systemic artery supply has been very well shown in a 41 mm embryo by Boyden. 25 These aberrant arteries are distinct from bronchial arteries, which are found to occur normally in these patients. 25

In most instances, in an area of lung with an abnormal arterial blood supply the tracheobronchial tree will be affected to some degree, so that there will be overlap with the following category. 32 33

In a fourth possibility (4 in fig 5), the insult is severe enough to disrupt both airway and pulmonary arterial development so that a systemic arterial supply is retained to the subsequent abnormal area of lung parenchyma. This is a combined bronchoarterial pulmonary malinosculation, and in its purest form would comply with most descriptions of "classical" sequestration.

The final possibility (5 in fig 5) is that, although there is considerable or complete disruption of the tracheobronchial connection, the pulmonary artery continues to develop and supply the abnormal lung segment. This will result in tracheopulmonary bronchopulmonary malinosculation with normal pulmonary artery blood supply, and include such entities as congenital lung cysts, so called sequestration with normal pulmonary artery supply, and cystic adenomatoid malformation. Peripheral lesions of terminal and respiratory bronchioles and alveoli (for example, congenital lobar emphysema) suggest a defect occurring late in development. Congenital lobar emphysema with systemic arterial blood supply has been described, supporting its embryological relationship to other forms of malinosculation.

This leaves extrapulmonary lung anomalies. In the early stages the developing lung bud, with its surrounding mesenchyme, lies in close proximity to the primitive foregut, from which it is derived. 24 28 Any insult or adhesion in an area where these two developing organs are adjacent is therefore likely to affect both structures; again, the timing and severity of the insult is important to the eventual outcome. Normally, the lung develops along genetically determined lines, the dividing epithelial cells being influenced by humoral factors from the surrounding mesenchyme. 10 34 Differentiating lung epithelial cells drawn by adhesions towards the primitive foregut may come to lie in mesenchyme, which is not conductive to their normal development. This would result in dysmorphogenesis of the misplaced tissue. 10 34

Primitive communications between this ectopic tissue and adjacent organ may develop at any stage. 10 34 28

Subsequent development of the pleura could sequestrate this tissue from its parent organ, either partially or completely, resulting in extrapulmonary sequestration or, alternatively, ectopic foregut derivatives within the lung. This hypothesis supports the contents of ourselves and others 3 6 8 13 27 that intrapulmonary and extrapulmonary lesions have a similar origin, without the need for an "accessory lung bud" theory. 28 It also explains the high incidence of associated congenital foregut anomalies occurring with extrapulmonary lung anomalies.

The development of the venous drainage is less well understood; but when anomalies of venous drainage occur in association with pulmonary malinosculation, they are likely to be secondary to the primary lesion and must be incorporated in the classification. Primary congenital anomalous venous drainage and anomalous systemic pulmonary artery collaterals secondary to congenital heart disease are not associated with any primary lung problem 35 36 and are therefore not included in the pulmonary malinosculation spectrum.

An interesting point is the absence of secondary...
parenchymal changes in patients with pulmonary systemic collaterals secondary to congenital heart disease (for example, in pulmonary atresia) even after many years. This confirms that the secondary changes that occur in malinosculation syndromes are not due to systemic pressures alone, but that there must be some underlying, inherent connective tissue abnormality within the lung neonatally.19 This is contrary to earlier hypotheses that parenchymal changes in malinosculation syndromes were primarily a systemic pressure effect.24

Summary

The pulmonary sequestration spectrum and related congenital lung anomalies present an extremely complex and varied group of bronchopulmonary-vascular malformations. Previous attempts at nomenclature and classification have proved inadequate. In this article we present a classification of the newly named pulmonary malinosculation spectrum, which includes all congenital lung anomalies where there is abnormal connection (that is, malinosculation) of one or more of the four major components of lung tissue—namely, tracheobronchial airway, lung parenchyma, arterial supply, and venous drainage, which in various combinations make up these lesions.

We feel that this simple descriptive approach will improve our understanding and management of these complicated lesions and this is supported by the clinical experience we report in the next article.

We would like to thank Professor Brian Corrin, Department of Pathology, for his helpful advice and criticism.

References


