Chest radiographic staging in allergic bronchopulmonary aspergillosis: relationship with immunological findings


Chest radiographic staging in allergic bronchopulmonary aspergillosis (ABPA) was addressed in 41 patients.

ASTRACT: The question of whether a chest radiographic severity staging system could be correlated with standard blood/serum diagnostic indices in allergic bronchopulmonary aspergillosis (ABPA) was addressed in 41 patients.

As asthma and positive Aspergillus fumigatus (AF) serology were considered essential diagnostic inclusion criteria. Eosinophil count, serum immunoglobulin (Ig)E and immediate skin hypersensitivity were also tested to grade patients as "definite" or "likely" ABPA. Definite cases had all five of these factors present, whereas likely cases had three or more. Chest radiographs were examined by experienced radiologists blinded to the clinical data. The six-stage radiographic score (0–5) was based on the severity and duration of changes seen: stage 0: normal; stage 1: transient hyperinflation; stage 2: transient minor changes; stage 3: transient major changes; stage 4: permanent minor changes; and stage 5: permanent major changes.

Significant positive correlations (p<0.05) were observed between peak AF titres (expressed as an index), peak eosinophil count and radiographic severity stage. When considered as subgroups, these correlations approached, but did not reach, significance for the group with "likely" ABPA (n=28), but in the group with definite ABPA (n=13), there was a high correlation between radiographic score and peak AF index (r=0.59), as well as peak eosinophil count (r=0.62).

This study suggests that the peak Aspergillus fumigatus index and eosinophil counts correlate best with the severity of radiographic stages in allergic bronchopulmonary aspergillosis. This chest radiographic staging system may be useful in the clinical assessment and management of patients with allergic bronchopulmonary aspergillosis, particularly in those patients with more severe radiographic stages.


Following the initial description of allergic bronchopulmonary aspergillosis (ABPA) by HOSKIN et al. [1] in 1952 in the UK, this condition has since been reported worldwide [2]. Fungi other than Aspergillus fumigatus (AF) have also been identified as causing a similar clinicopathological process [3, 4]. HENDERSON et al. [5], in a study of 107 consecutive patients admitted to a hospital for chronic chest disease, reported 22% of 46 asthmatics to have ABPA. Another study indicated that the prevalence period >4 yrs of allergic bronchopulmonary mycosis in a hospital outpatient respiratory medicine department was just >1% [6]. There is still an incomplete agreement as to what clinical, radiographic, and laboratory features are necessary to diagnose ABPA, but ROSENBERG et al. [7] and SAKSIN et al. [8] are in broad agreement and recommend six of the following “primary criteria” for definite diagnosis: episodic bronchial obstruction (asthma), peripheral blood eosinophilia, immediate skin reactivity to aspergillus antigen, precipitating antibodies against aspergillus antigen, elevated serum immunoglobulin (Ig)E, history of pulmonary infiltrates (transient or fixed) and central bronchiectasis. In pulmonary sarcoidosis, chest radiographic severity staging (stages 1–4) has been used as an index of activity of this disease and of prognosis [9]. A clinical staging system has been suggested for use in ABPA [10]. The question was addressed as to whether a chest radiographic staging system in ABPA could be related to blood/serum diagnostic criteria, i.e. eosinophil count, serum IgE and aspergillus antibodies. If a correlation to these criteria was found it was proposed that chest radiographic staging may be a useful monitor of disease activity and possibly of prognosis, analogous to the use of chest radiographic staging in pulmonary sarcoidosis.

Patients and methods

Patients selected for inclusion in this study attended the Respiratory Medicine clinic at the University Hospital (Cork, Ireland) during the 11 yrs 1985–1995. There was a diagnosis of asthma in all cases. Patient details are listed in table 1 and all five of the nonradiological criteria proposed by ROSENBERG et al. [7] and SAKSIN et al. [8] were used...
Dublin, Ireland). Internal positive controls (positively react-
in the secondary reaction. Tests were performed using a
were used in this process. Alkaline phosphatase-conj-
immunosorbent assay (ELISA). Somatic and culture fil-
tasis. The highest grade reported was used for correlation
indicates <6 months duration. Grade 4 was identified by
hyperinflation without other changes, Grade 2 reflected
a normal radiograph, Grade 1 was assigned to transient
marginal changes such as "gloved finger" or band
changes such as "fibrosing alveolitis", lobar shrinkage and
consolidation, parallel line and ring shadows, nodules,
avascular areas, "honeycombing", "toothpaste" shadows,
globed finger", band shadows and tramline shadows.
Changes like "fibrosing alveolitis", lobar shrinkage and
atelectasis, as well as pseudohilar adenopathy and pleural
thickening have also been described [11–16]. No radio-
herapy, hyperinflation, various infiltrate patterns,
consolidation, parallel line and ring shadows, nodules, avascular areas, "honeycombing", "toothpaste" shadows,
changes such as "fibrosing alveolitis", lobar shrinkage and
atelectasis, lobar shrinkage or proximal bronchiolec-
tasis. The highest grade reported was used for correlation
with blood/serological data.
Antibodies to AF were measured by enzyme-linked
immunosorbent assay (ELISA). Somatic and culture fil-
trate antigens of AF (Mercia Diagnostics, Dublin, Ireland)
were used in this process. Alkaline phosphatase-conju-
gated antihuman IgG (Boehringer, Dublin, Ireland) and
p-nitrophenylphosphate (Sigma, Dorset, UK) were used
in the secondary reaction. Tests were performed using a
microtitre reader (Minireader MR 590; Shaw Scientific,
Dublin, Ireland). Internal positive controls (positively reac-
ting sera) were standardized against commercially obtained
positive controls (Mercia Diagnostics).
Results of peak AF antibody titres were expressed as an
index or ratio. Positively reacting sera gave readings that
were at least 2.5 times that of negative controls. Peak eosino-
phils were also used. Total serum IgE was mea-
sured by a solid-phase immunoassay (Unicap Allergy
System, Pharmacia, Dublin, Ireland).

### Statistical analysis

The data for chest radiographs grade, peak aspergil-
lus indices and peak eosinophil counts were not normally
distributed; therefore, correlations were determined using
Spearman’s rank order method. Analysis of total serum
IgE data included log transformation. Statistical signifi-
cance was determined by p-values <0.05.

#### Results

Significant correlations were found between peak asp-
erillus index and chest radiographic stage and between
peak eosinophil count and chest radiographic stage among
the group as a whole (table 2 and fig. 1a) and these same
correlations were higher for the group considered to have
"definite" ABPA when considered as a separate subgroup
(table 2 and fig. 1). Correlations for the same parameters
tended towards, but did not reach significance in the group
with "likely" ABPA (table 2 and fig. 1a). Total serum IgE
did not correlate with chest radiographic stage.

#### Discussion

A wide variety of described chest radiographic changes
are associated with ABPA. These changes include normal
radiography, hyperinflation, various infiltrate patterns,
consolidation, parallel line and ring shadows, nodules, avascular areas, "honeycombing", "toothpaste" shadows,
gloved finger", band shadows and tramline shadows.
Changes like "fibrosing alveolitis", lobar shrinkage and
atelectasis, as well as pseudohilar adenopathy and pleural
thickening have also been described [11–16]. No radio-
graphic staging system has been widely accepted for use
in this disease. Previous data failed to show any correla-
tion between the immunological findings on the one hand
and clinical findings, such as age, duration of asthma or
aspergillosis, but were able to demonstrate significant

### Table 1. – Patient characteristics

<table>
<thead>
<tr>
<th>Age yrs</th>
<th>Aspergillus index</th>
<th>IgE level IU·mL⁻¹</th>
<th>Eosinophil count cells·µL⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Whole group (n=41)</td>
<td>47.71</td>
<td>4.15</td>
<td>820.00</td>
</tr>
<tr>
<td>&quot;Definite&quot;</td>
<td>(19.46)</td>
<td>3.4</td>
<td>1000</td>
</tr>
<tr>
<td>ABPA (n=13)</td>
<td>47.92</td>
<td>5.05</td>
<td>798.50</td>
</tr>
<tr>
<td>&quot;Likely&quot; ABPA (n=28)</td>
<td>45.16</td>
<td>6.05</td>
<td>798.50</td>
</tr>
</tbody>
</table>

Age is given as mean (±); other data are presented as median values. Laboratory reference range for Aspergillus fumigatus
index: positive >2.5; reference range for immunoglobulin (Ig)E upper limit of normal=150 IU·mL⁻¹; upper limit for eosino-
phils=450 cells·µL⁻¹. ABPA: allergic bronchopulmonary asper-
gillosis.

in patient selection. All patients had positive AF serology.
Cases negative for AF serology were excluded. The pres-
ence of peripheral blood eosinophilia, positive immediate
skin reactivity to prick testing with AF antigen and ele-

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in this disease. Previous data failed to show any correla-
tion between the immunological findings on the one hand
and clinical findings, such as age, duration of asthma or
aspergillosis, but were able to demonstrate significant

### Table 2. – Correlation of peak aspergillus indices and
eosinophil counts with chest radiographic grade

<table>
<thead>
<tr>
<th>Aspergillus index</th>
<th>Eosinophil count</th>
</tr>
</thead>
<tbody>
<tr>
<td>r</td>
<td>p</td>
</tr>
<tr>
<td>Whole group (n=41)</td>
<td>0.39</td>
</tr>
<tr>
<td>&quot;Definite&quot; ABPA (n=13)</td>
<td>0.59</td>
</tr>
<tr>
<td>&quot;Likely&quot; ABPA (n=28)</td>
<td>0.33</td>
</tr>
</tbody>
</table>

ABPA: allergic bronchopulmonary aspergillosis.
relationships between the aspergillus precipitin tests and the radiologically assessed activity of the disease. In the same study [17] IgE levels were not found to have a linear correlation with radiographic changes. These data support the present findings and suggest that AF antibody levels are more closely related than IgE to changes in chest radiographic images.

One would expect higher aspergillus indices and higher degrees of eosinophilia during periods of increased symptomatology or flare-ups to coincide with temporary changes on chest radiography. However, as the current methodology disassociated the highest grades on chest radiographs from the blood/serum indices, the relationship between these variables may be due to the reasoning that, over time, patients with flare-ups will eventually progress to some degree of pulmonary scarring. This may appear on the chest radiograph and therefore cause this association.

The present data indicate that the group with "definite" ABPA by our criteria have a significant correlation between both the AF index and chest radiographic stage, and between eosinophil count and chest radiographic stage. The trend in the group with "likely" ABPA, although not reaching statistical significance, is generally in keeping with that of the "definite" group and supports the link.

Fig. 1. – Correlations of a) aspergillus index and b) eosinophil count with chest radiographic grade in the whole group. Correlations of c) aspergillus index and d) eosinophil count with chest radiographic grade in "definite" allergic bronchopulmonary aspergillosis (ABPA). Correlations of e) aspergillus index and f) eosinophil count with chest radiographic grade in "likely" ABPA.
between the severity of radiographic changes and higher AF indices and eosinophil counts.

A chest radiographic staging system such as the above, together with peak Aspergillus fumigatus indices and eosinophil counts, may be useful in delineating the activity of the disease and, in addition, may help to predict the likelihood of progression of the disease. Accordingly, this staging system may also help in the clinical assessment and management of allergic bronchopulmonary aspergillosis, particularly in those patients with more severe chest radiographic stages.

References