Diagnostic approach in cases with suspected work-related asthma

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Abstract

Background: Work-related asthma (WRA) is a major cause of respiratory disease in modern societies. The diagnosis and consequently an opportunity for prevention are often missed in practice.

Methods: Based on recent studies and systematic reviews of the literature methods for detection of WRA and identification of specific causes of allergic WRA are discussed.

Results and Conclusions: All workers should be asked whether symptoms improve on days away from work or on holidays. Positive answers should lead to further investigation. Spirometry and non-specific bronchial responsiveness should be measured, but carefully performed and validly analysed serial peak expiratory flow or forced expiratory volume in one second (FEV1) measurements are more specific and confirm occupational asthma in about 82% of those still exposed to the causative agent. Skin prick testing or specific immunoglobulin E assays are useful to document allergy to high molecular weight allergens. Specific inhalational challenge tests come closest to a gold standard test, but lack standardisation, availability and sensitivity. Supervised workplace challenges can be used when specific challenges are unavailable or the results non-diagnostic, but methodology lacks standardisation. Finally, if the diagnosis remains unclear a follow-up with serial measurements of FEV1 and non-specific bronchial hyperresponsiveness should detect those likely to develop permanent impairment from their occupational exposures.

Keywords: Asthma, Occupational disease, Diagnosis, Work-related

Background

At least 15% of adult asthma is induced or triggered by factors in the workplace [1]. Early diagnosis can improve the prognosis of work-related asthma since cessation of exposure after appearance of asthmatic symptoms and identification of specific sensitization within the first months after onset of symptoms, may permit a full recovery. In addition, diagnosis of a case of work-related asthma (WRA) is a sentinel event that should lead to effective primary preventive measures in the individual workplace or in a branch of industry [2-4]. The reader is referred to recent comprehensive and systematic literature reviews [5-11]. Diagnosis of work-related asthma is in many cases a complex undertaking consisting of various diagnostic tests and procedures. The main object of diagnosis is to identify the cause of the asthma, in particular to separate occupational asthma from asthma unrelated to work, assisting decisions about intervention to prevent further exposure to the patient and is also important for handling compensation claims. The possibilities for prevention and degree of economic support for the patients may differ between industries and countries. The optimal diagnostic strategy will vary depending on medical as well as economic and juridical evidence.

Obvious preconditions for considering a diagnosis of WRA are typical diagnostic findings of asthma (see below) and an association of asthma symptoms with exposure at work. Asthma present before occupational exposure but associated with worsening at the start of a new occupational exposure indicates a diagnosis of work-aggravated asthma (WAA). Asthma relapsing or becoming more severe some time after exposure to a new occupational allergen may be due to occupational asthma, and does not...
preclude the diagnosis. A thorough clinical and occupational history should be obtained in all cases [12,13].

The diagnosis of WRA can be performed in three steps:

1. Making the diagnosis of asthma.
   Asthma diagnosis is performed according to recent international guidelines [14,15]. However, the diagnosis of asthma is not always straightforward. Asthma is a heterogeneous disease, and the diagnostic criteria are ill-defined and vary between studies [16,17]. Further, there is considerable overlap between asthma and chronic obstructive pulmonary disease (COPD) [18]. A doctor’s diagnosis of asthma may be frequently questionable [19].

2. Identification of the workplace as the cause of the patient’s asthma.
   Objective verification of the association between work exposure and airflow limitation is a basis for diagnosing WRA, both for new onset occupational asthma (OA) and WAA.

3. Identification of a specific agent causing WRA.
   This step is most demanding on resources. A specific diagnosis is needed before appropriate remedial action can be taken in the workplace, but the full diagnostic armamentarium is only available at few centres worldwide [20]. The extent of diagnostic investigation will depend on what resources are available and on a decision analysis that takes into account the consequences of false positive and false negative diagnoses [21].

**Diagnosis of allergic work-related asthma**

Recognized exposure to a known allergen at work and a provisional diagnosis of WRA should lead to an extensive diagnostic work-up to identify the agent and objectively confirm its causal role. Occupational allergens conform to the general definition of allergens [22].

To reach the diagnosis of WRA several pieces of evidence are necessary:

- Objective information of exposure to known sensitizers.
- Demonstration of an association between exposure at work and airflow limitation by lung function testing or changes in non-specific bronchial hyperresponsiveness (NSBHR) or changes of airway inflammation (e.g. sputum eosinophils).
- Identification of a specific allergic reaction to the occupational agent (rather than an acute irritant reaction).

Note: There are many OA-inducing agents and worksites generally included in the lists of airway allergens and irritants (for details see latest summaries [23,24] and Irritants: Evidence table of reporting OA or occupational COPD due to irritants (Authors: X. Baur, P. Bakehe, H. Vellguth, JOMT), ACGIH Tables: Occupational agents with respiratory effects according to ACGIH and/or classified with the H334 (R42) phrase and/or H335 (R37) phrase plus EU regulations (Author: X. Baur); however, due to negative allergological findings for many of them pathogenetic mechanisms are still unknown; examples include most cases of isocyanate and potroom asthma [25].

Diagnostic tests can be divided into those which should be available to all respiratory physicians and those generally confined to specialist in occupational lung diseases.

**Diagnostic methods suitable for all respiratory and occupational physicians**

All working patients with asthma and COPD should be asked whether their symptoms improve on days away from work or on holidays. Further investigation is required for all positive respondents.

**Questionnaires.** Questionnaires are extensively used as screening tools in epidemiological studies. Several epidemiological questionnaires are in current use [26-28]. Questions about work exposure and association of symptoms to work may be used in clinical diagnosis. In the clinical setting questionnaires that identify symptoms of wheeze and/or shortness of breath which improve on days away from work or on holiday have a high sensitivity, but relatively low specificity for OA [28-34].

**Medical histories by experts.** There is general agreement that medical histories taken by experts have high sensitivity, but their specificity may be lower [28,29,32,35-38].

**Exposure assessment** [39]. This comes both within the compass of a generalist and a specialist. A generalist should elicit exposures in high-risk occupations, in which asthma should be assumed to be occupational unless excluded by objective tests. *The workers reported from population studies to be at increased risk of developing asthma include; bakers, chemical workers, cleaners, cooks, electrical and electronic production workers, farm workers, food processors, forest workers, healthcare workers, laboratory technicians, mechanics, metal workers, painters, plastics and rubber workers, storage workers, textile workers, waiters, welders and wood workers. A specialist should identify specific exposures, with a systematic occupational history, scrutiny of exposure documentation such as material safety data sheets (MSDSs), internal reports and industrial hygiene measurements from the industry. The failure to find a sensitiser on an MSDS should not preclude the diagnosis of occupational asthma, as many sensitisers are not regularly listed, particularly those in low concentration, those that are only...*
present in certain circumstances, such as when heated and those given non-specific titles such as preservative, biocide, fragrance, resin etc.

**Spirometry** [40,41]. Spirometry is required in all patients considered for WRA. It is used as a main instrument for monitoring lung function longitudinally during surveillance, also during measurement of non-specific bronchial hyperresponsiveness (NSBHR) and in specific inhalation challenge. Many workers with WRA have normal spirometry when seen in a clinic. There is also a role for measurement of airway resistance or specific conductance to monitor lung function when a patient cannot record FEV₁ reliably.

**Pre- to post-shift changes in lung function** cannot be recommended for the validation or exclusion of WRA [42,43]. Serial PEF and spirometry measurements have been shown to be superior to cross-shift change in diagnosing WRA [43].

**Measurement of non-specific airway responsiveness** [44,45]. Challenge with methacholine is part of the initial diagnostic work-up. It can also be measured before and after a period of work exposure and during SIC. Normal NSBHR does not exclude OA. A large number of concordant studies from different centres using different methodologies demonstrated that increased NSBHR is often found in workers with WRA. There are, however, many reports of normal methacholine or histamine reactivity within 24 hours of exposure in workers with confirmed OA [28,29,32,34,36,38,46-57].

**Serial lung function testing.** If clinical OA exists, exposure to routine levels of the causative agent should result in a measureable decline in lung function in the hours after exposure. This is usually performed by a simple expiratory peak flow (PEF) meter or by a portable spirometer [58]. Measuring PEF or FEV₁ during periods of work and away from work may document a causal relationship between exposure and airflow reduction [58]. Technical issues related to interpretation have been extensively studied [59]. The frequency and duration of the recording depends on the method of analysis, but in general, at least 4 readings/day are required, although 8 readings a day allows shorter records, with usual exposure on work days and no changes in treatment on days away from work. A 3-week record containing at least 3 periods off work would be suitable for most forms of analysis [60-62]. The sensitivity and specificity of serial peak flow measurements performed after extensive instruction to patients and using quality control are high in the diagnosis of OA, for which there is a recent systematic review with a pooled sensitivity of 82% and specificity of 88%. Peak flow measurements require encouragement and instruction of the patient; otherwise at least one third of the subjects will not produce reliable data [30,31,43,48,56,62-68]. Note: New handheld electronic spirometers storing the whole flow volume curves and automatically checking for reproducibility and other quality criteria (e.g. ATS/ERS 2005) will allow easy quality control and provide additional parameters [41]. Their practicability, including the analysis of the large data set that can result from testing one patient, remains to be shown.

**Allergological tests**

- The skin prick test (SPT) is the mainstay of allergological testing. SPT is generally regarded as sensitive, but with less specificity. The test should be performed according to international guidelines [69]. Standardized material is rarely available in suspected occupational allergy cases, but crude extracts produced locally may be useful. SPT is usually not performed in low molecular weight (LMW) allergy (with the exception of platinum salts and reactive dyes).

- Specific immunoglobulin E (IgE) tests are often less sensitive, but possibly more specific than SPT. A major shortcoming of allergological tests is the lack of standardization of most occupational allergens as well as of the antibody measuring assays [70-72]. Both SPT and specific IgE tests are in general sufficiently sensitive for detecting type I sensitisation and OA caused by most high molecular weight agents (HMW), but are not specific for diagnosing asthma [34,73]. Both SPT and specific IgE tests are sensitive for detecting type I sensitisation and occupational asthma caused by acid anhydrides and some reactive dyes, but have a lower specificity for diagnosing asthma in these cases [74-78]. Skin prick tests are highly sensitive but less specific for occupational asthma caused by complex platinum salts [53,54].

- Other in vitro studies such as mediator release test (i.e. of histamine) in an allergen-antibody reaction and release of cytokines during in vitro stimulation [79] are, to date, not routine tests.

**Diagnostic methods suitable for specialists in occupational lung diseases**

Specific inhalation challenge tests (SIC) [80,81]. SIC are commonly regarded as a reference method for diagnosing sensitizer-induced WRA. However, the test is in practice not well standardized internationally and is sparsely available [82]. They are subject of a current ERS taskforce whose preliminary indications are:

- Confirming the diagnosis and cause of OA when other objective methods are not feasible, have failed or provide indeterminate results.
As a rapid way to evaluate the possibility of OA caused by a sensitizing agent when the net potential clinical benefit of the test is expected to exceed the risk of complications.

When a new (not formerly described) specific cause of OA is suspected.

For research purposes.

Methods for performance of specific inhalation tests:
These may be performed in a specially designed laboratory in two main ways:

1. Controlled laboratory challenge.
   By use of this method controlled concentrations of the suspected sensitizers (standardized allergen extract) are administered in controlled concentration and dose either by the inhalation chamber method or by closed-circuit methodology [81]. Laboratory-based specific challenge testing may produce false negative results, which has been shown when comparing laboratory challenge with workplace challenges which confirmed occupational asthma [83].

2. Workplace challenge test or realistic challenge.
   This is an attempt to reproduce actual work processes and exposures which are performed in line with the Jack Pepys tradition [84].

The aim of a workplace challenge is to make supervised lung function similar to a bronchial provocation testing in the workplace after suitable control measurements have been made without exposure in the workplace on separate days. Measurements of NSBHR, cells in induced sputum, fractional exhaled nitric oxide (FeNO) etc. can be made in conjunction with the workplace challenge. The patient should be exposed to the usual concentrations of the potential causative agent during the challenge that may last up to 2−4 hours of usual exposure. It is important to ensure that the usual work practices are taking place during the workplace challenge which may not always be easy to achieve.

Safety considerations are important, and many workplaces do not provide clean and safe environments for medical treatment and the whole site may be the cause of the asthma. In these circumstances any emergency treatment of a severe asthmatic attack would involve removing the worker from the workplace and often from sources of electrical supply, typically needed to perform spirometry and operate nebulizers.

Limitations of specific inhalation challenge
Specific bronchial provocation testing can produce false positive and false negative results. It is particularly difficult to set up when the exposures are complex (such as with welding fume or metal working fluid exposures) or when the process being mimicked involves high temperatures, explosive or inflammable material. SIC is in general poorly standardized. Even if the procedure is usually regarded as the gold standard for diagnosing WRA, it requires much experience to adequately define the challenge procedure and to know which parameter to use when assessing the sensitivity and specificity of this test.

The earliest attempt at assessing the validity of SIC used continued work with exposure to the original suspected causative agent without reduction of exposure for a year. The subsequent absence of deterioration in asthma was taken as an indicator of no WRA, and the need for removal from exposure because of repeated severe asthmatic attacks indicated OA. Using these two criteria, the sensitivity for specific bronchial provocation testing with isocyanates up to a 30 minute exposure to 0.02 ppm was 82% and for colophony up to a 15 minute exposure to heating neat colophony was 100%. The relevant specificities were 67% for colophony and 100% for isocyanate exposure [85].

More recently Rioux [83] performed work place challenges in 99 workers who had negative specific bronchial provocation testing but had a good history of OA. Twenty-nine of these had a positive workplace challenge, some with subsequent positive bronchial provocation testing when different agents were tested. Of those who had negative workplace challenges, 34/70 had asthma or rhinitis excluded following further investigation. False negative specific challenges were therefore found in 29/65 with asthma or rhinitis and a good history of workplace deterioration [83].

There are no studies evaluating the criteria for a positive or negative workplace challenge. However, the criteria for detecting late asthmatic reactions from specific challenges proposed by Stenton and colleagues [86] are probably appropriate. These criteria specify at least three control days without exposure, calculating the pooled standard deviation for the FEV1 measurements on the unexposed days and requiring at least 2 consecutive measurements on the workplace challenge day to be below these pooled standard deviation measurements [86]. This method has been applied to unsupervised serial measurement of PEF and found to have a specificity of over 90% with specific challenge testing as the standard [87].

In summary, in spite of their limitations carefully controlled specific challenges come closest to a gold standard test for some agents causing OA [42,43,49,52]. Further, a negative test in a worker with otherwise good evidence of OA is not sufficient to preclude the diagnosis [42,43,49,52].

Non-invasive methods to assess airway inflammation
Several non invasive methods have been used to assess and monitor airway inflammation in asthma in the last decade.
Three of these methods have been standardized by international committees:

- induced sputum
- exhaled nitric oxide (FeNO)
- exhaled breath condensate (EBC).

Analysis of induced sputum is a valid and reproducible method for studying airway inflammation [88,89]. The method consists of inducing sputum production by having the patient inhale a hypertonic saline solution. Sputum is subsequently processed and cytospins are prepared and stained. In addition, inflammatory mediators can be measured in sputum supernatant [89]. The count of eosinophils is the most repeatable parameter in induced sputum and is a reasonably good non-invasive index of airway eosinophilia since there is fairly good agreement between the evaluations of eosinophils in induced sputum, in bronchial biopsy specimens, and in bronchoalveolar lavage [90]. The validity of this method is attested to by several studies, which have shown that sputum inflammatory indices such as eosinophils and eosinophil cationic protein (ECP) are increased by exposure to common allergens and are reduced with corticosteroids [91]. There is evidence that induced sputum may be useful in clinical practice since evaluation of sputum eosinophils was shown to be equivalent to NSBHR for discriminating asthma from other conditions, and assessment of sputum eosinophilia improved the management of asthma [17,92,93].

Several studies reported sputum eosinophilia in both early and late reactors between 6 and 24 hours after SIC in OA caused by both high- and low-molecular weight occupational agents. Moreover, the addition of sputum cell counts to monitoring of PEF increased the specificity of this test suggesting that monitoring airway eosinophils in induced sputum improves the diagnosis of OA [94-96]. Indeed, induced sputum may be useful in the diagnosis and follow-up of subjects with WRA [97]. Its utility in epidemiological studies has not been evaluated.

Eosinophil counts in sputum may have other diagnostic uses. Sputum might be helpful in differentiation between WAA and OA due to a workplace sensitiser that is superimposed on existing asthma, since two studies have shown that exposure to occupational agents in asthmatics not sensitised to the agents did not induce airway inflammation and did not change the sputum cell composition [88,98]. The analysis of induced sputum has also been found to be useful in the identification of occupational eosinophilic bronchitis. This condition is characterized by cough on exposure to occupational agents, without any change in lung function tests [99].

Sputum eosinophilia is not present in all workers with allergic WRA. Compared with those with sputum eosinophilia, those without eosinophilia were exposed to similar agents, had similar changes in lung function related to work exposure but had less NSBHR [46].

Inhalation of hypertonic saline is not completely non-invasive, and can cause bronchoconstriction in subjects with hyperresponsive airways. Its success rate is less than 100%, since some subjects are unable to produce adequate samples. In addition, the induction is time-consuming and processing the samples is laborious and needs to be performed soon after collection [89]. The conclusions from the ERS evidence-based guidelines was [10];

- Sputum eosinophils increasing by >1% post SIC or workplace exposure may support a diagnosis of OA when the FEV1 has fallen <20%.
- The presence or absence of increased sputum eosinophils is not useful in selecting or excluding those who might have work-related asthma.

Exhaled nitric oxide (FeNO) is a non-invasive tool for assessing airway inflammation in asthma [100]. NO is produced by various cells in the lung either resident or recruited during the inflammatory process. Exhaled NO is generally measured on line, the subject blowing directly into the analyser with immediate results. Portable analysers are also available. It is also possible to have a remote breath collection into inert bags, with subsequent analysis. NO concentration is increased in the exhaled air from patients with asthma and decreased by corticosteroid therapy [101-103]. There is a positive correlation between FeNO and sputum eosinophils in asthma [104]. In clinical practice, evaluation of FeNO was equivalent to sputum eosinophils in diagnosing asthma [105] and management of asthma according to FeNO levels was effective to reduce maintenance doses of inhaled corticosteroids [106].

Some occupational studies have investigated the role of FeNO in assessing OA, but with inconsistent results [98,102,107-112]. It has been suggested that measurement of FeNO can be used to indicate the development of airway inflammation accompanying late asthmatic reactions after SIC in patients with normal or slightly increased basal NO levels [112]. However, the usefulness of FeNO in the investigation of OA may be limited by factors affecting its determination, such as therapy with inhaled steroids and smoking [100]. Although the measurement of FeNO is totally non-invasive, quick and relatively simple to perform, more data are needed to determine whether FeNO is useful in diagnosing WRA.

- In the clinical setting, a finding of normal exhaled nitric oxide fraction cannot be used to exclude OA.

EBC is collected by cooling or freezing exhaled air and is also totally non-invasive [113,114]. In addition to
measuring the pH of EBC, investigators have been able to detect several volatile and non-volatile chemicals, including lipids, proteins and oxidants. Nevertheless, there are still open questions in the procedure to collect samples, the reliability of analytical methods for detecting minute concentrations of chemicals and the expression of the results. EBC has been used in the investigation of airway diseases such as asthma, COPD, lung cancer, interstitial lung disease and acute respiratory distress syndrome. Since no studies using EBC have been performed in OA so far, its role in the diagnosis of WRA remains undetermined.

In summary, the above-mentioned new methods are promising for monitoring the inflammatory activity in the airways. However, their definite place in practical diagnosis is at present not clear.

**Diagnosis of diagnostic groups within the spectrum of WRA**

Individual diagnostic steps of the following WRA forms correspond to the methods mentioned earlier in the section on diagnosis of allergic WRA. However, SIC may produce positive results in spite of negative IgE findings, e.g. in cases suffering from isocyanate, platinum or irritant asthma with latency (it is clearly inappropriate for acute irritant induced asthma).

1. Work-aggravated asthma. This is an entity characterized by symptomatic deterioration of pre-existing or concomitant non-occupational asthma at work without a latent interval mainly by irritant mechanisms. There are similar changes in lung function related to work exposure as those with allergen- or irritant-induced OA [115].
2. Irritant-induced asthma [116].
   a. Acute irritant induced asthma (Reactive airways disease syndrome, RADS) [117]. This is a fairly well described entity. The diagnosis is retrospective and criteria based, e.g. ACCP criteria for diagnosis of reactive airway dysfunction syndrome (RADS) (should meet all eight) [12]:
      1. Documented absence of preceding respiratory complaints.
      2. Onset of symptoms after a single exposure incident or accident.
      3. Exposure to a gas, smoke, fume, or vapor with irritant properties present in very high concentration.
      4. Onset of symptoms within 24 h after exposure with persistence of symptoms for at least three months.
      5. Symptoms simulated asthma: cough, wheeze, dyspnoea.
6. Presence of airflow obstruction on pulmonary function tests (initial testing should be done shortly after exposure).
7. Presence of nonspecific bronchial hyperresponsiveness.
8. Other pulmonary diseases ruled out.
b. Not so sudden onset of irritant asthma. As opposed to RADS this is caused to lower chronic or repetitive exposures to irritants mostly in the range of the occupational exposure limits [8,118], it is commoner in those with atopy or childhood asthma.
c. Irritant asthma with latency [119]. This is a controversial entity of asthma defined by
   - no prior asthma
   - a latent interval from first exposure to disease
   - no massive acute exposure
   - symptoms related to usual exposure to the causative agent
   - reproducibility of the asthma from either workplace challenges or specific inhalation challenge or valid serial measurements of PEF measurements at and away from work
   - an allergic mechanism is very unlikely

**Discussion**

The problem of test materials and lack of standardized technology:

Immunological tests are commonly performed as used in allergy testing, but the main problem in occupational allergy is an almost universal lack of standardized requirements for allergy extracts. Simple test may be performed using material that patients bring to the consultation. For HMW agents close collaboration with an immunochemist is advisable for optimal characterization of test material. As regards LMW agents, the problem is even more complex. For instance, there are more than thousands different isocyanates compounds known as yet. In research, most publications deal with monomers such as toluene disiocyanate, hexamethylenediisocyanate, methylene diphenyldiisocyanate. However, some patients may react only with oligomers that are predominantly used by the industry [120,121]. Furthermore, a new generation of isocyanates results from the decomposition of polyurethane coatings or foam during hot work, a process that can yield irritant toxic agents as well [122]. Collaboration with a qualified chemist is thus necessary to plan and monitor tests, especially SIC with complex chemicals.

**Situations with conflicting test results**

WRA diagnostic efforts may provide conflicting findings, e.g. the patient may report convincingly work-related asthmatic symptoms but there are no abnormal findings
in lung function and allergological tests. Follow-up of the subject at short intervals including serial FEV1 and NSBHR monitoring is recommended in such situations. An accelerated FEV1 decline can be detected statistically using the Spirola program [123].

Conclusions

The potential health benefit of a correct and early diagnosis of WRA is considerable both for the individual, industry and society. However, the possibility of WRA is frequently not considered by the doctor and adequate measures to secure an etiologic diagnosis are not performed. Organized collaboration between occupational, primary care and specialized centre physicians is required to improve the situation. An integrated approach is recommended [71].

Abbreviations

NSBHR: Non-specific bronchial hyperresponsiveness; WRA: Work-related asthma; OA: Occupational asthma; PEF: Peak expiratory flow; SPT: Skin prick test; SIC: Specific inhalation challenge.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

TBAa drafted the manuscript. All co-authors made substantial contributions to the review. All authors read and approved the final manuscript.

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