
There may be differences between this version and the published version. You are advised to consult the publisher’s version if you wish to cite from it.

http://eprints.gla.ac.uk/120917/

Deposited on: 15 July 2016
### Addressing corticosteroid insensitivity in adults with asthma

<table>
<thead>
<tr>
<th>Journal:</th>
<th>Expert Review of Respiratory Medicine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manuscript ID</td>
<td>ERRX-2015-0097.R1</td>
</tr>
<tr>
<td>Manuscript Type:</td>
<td>Reviews</td>
</tr>
<tr>
<td>Keywords:</td>
<td>Adherence, Asthma, Biologic agents, Biomarkers, Cigarette smoking, Corticosteroids, Corticosteroid insensitivity, Corticosteroid resistance, Small molecule drugs</td>
</tr>
</tbody>
</table>
Title: Addressing corticosteroid insensitivity in adults with asthma

Abstract

Corticosteroids are the most effective treatment for asthma, but the therapeutic response varies markedly between individuals, with up to one third of patients showing evidence of insensitivity to corticosteroids. This article summarizes information on genetic, environmental and asthma-related factors as well as demographic and pharmacokinetic variables associated with corticosteroid insensitivity in asthma. Molecular mechanisms proposed to explain corticosteroid insensitivity are reviewed including alterations in glucocorticoid receptor subtype, binding and nuclear translocation, increased proinflammatory transcription factors and defective histone acetylation. Current therapies and future interventions that may restore corticosteroid sensitivity in asthma are discussed, including small molecule drugs and biological agents. In the future, biomarkers may be used in the clinic to predict corticosteroid sensitivity in patients with poorly controlled asthma.

Word count: 120 words

Key words: Adherence; asthma; biological agents; biomarkers; cigarette smoking; corticosteroids; corticosteroid insensitivity; corticosteroid resistance; small molecule drugs.
1. Introduction

Asthma is a chronic inflammatory disease of the airways that affects 300 million people worldwide. Both national and international guidelines recommend daily inhaled corticosteroid as the preferred controller treatment for adults and adolescents with asthma who have poor symptom control or who at risk of exacerbations [201, 202]. Inhaled corticosteroid use in asthma reduces symptoms, improves quality of life and increases lung function as well as decreases the rate of exacerbations [201, 202]. The majority of the therapeutic benefits of inhaled corticosteroids are achieved at low to medium doses [1], although higher doses are often required in patients with more severe asthma [2] [202]. Short courses of oral corticosteroids are administered to treat severe exacerbations and daily oral corticosteroids are used in the lowest dose providing adequate control to treat patients with severe asthma when symptoms are uncontrolled despite maximal therapy [201, 202].

Guideline recommendations for inhaled and oral corticosteroid use in asthma are based on average therapeutic responses from clinical trial populations that may not be representative of ‘real-life’ patients [3]. Numerous studies in adults and children with asthma have noted a considerable patient-to-patient variability in improvements in lung function and airway hyperreactivity with inhaled and oral corticosteroid treatment in patients with apparently similar levels of disease severity [4-6] (Figure 1). The findings from these studies suggest that corticosteroid insensitivity may be present in approximately one third of patients with asthma. Inhaled corticosteroids exhibit a linear dose-response for systemic adverse effects [7] and high doses are associated with adrenal suppression [1,8], reduced bone mineral density and increased risk of fractures as well as diabetes and reactivation of tuberculosis [9]. To improve the
therapeutic ratio for corticosteroid use in asthmatic patients who have an impaired clinical
responses to corticosteroids there is a need to understand the causes of corticosteroid
insensitivity and how best to manage these patients. This article reviews published information
on factors associated with corticosteroid insensitivity in asthma, considers the molecular
mechanisms proposed to explain corticosteroid insensitivity and discusses current therapies and
future interventions that may restore corticosteroid sensitivity in asthma.

2. Factors associated with corticosteroid insensitivity in asthma

Corticosteroid insensitivity is an imprecise term used to describe an impaired response to
corticosteroids, whereas corticosteroid resistance implies the absence of any response to
corticosteroids. A range of in vivo and in vitro biomarkers have been used to assess the
response to corticosteroids in man, although none are ideal. The variation in corticosteroid
sensitivity in people with asthma is most commonly measured using clinical outcomes such as
symptoms, lung function and exacerbations [6]. A composite phenotype that includes
information on inhaled corticosteroid treatment response in patients with mild to moderate
asthma assessed by six outcomes (lung function, bronchodilator response, airway
responsiveness, symptoms, need for oral corticosteroids and frequency of emergency
department visits and hospitalizations) is reported to identify endophenotypes that more
accurately describe corticosteroid sensitivity [10]. Sputum eosinophils, transcription factor
gene expression levels on airway cells [11] and exhaled nitric oxide [12] as well as bronchial
allergen challenge [13], exhaled breath profiling [14], urinary biomarkers [12] and the
cutaneous vasoconstrictor response to topical corticosteroid [15] have been employed to
measure variability in corticosteroid sensitivity in asthma. In vitro tests of corticosteroid
induced suppression of peripheral blood mononuclear cell (PBMC) T lymphocyte proliferation [16] and expression of transcription factors on cells [16,17] or composites in vitro measurements [18] have been used as biomarkers of corticosteroid sensitivity, although the ability of these tests to predict clinical responses to corticosteroids in the general population of patients with asthma is uncertain.

Genetic, environmental and asthma associate factors as well as demographic and pharmacokinetic variables may contribute to differences in sensitivity to corticosteroids between people with asthma (Table 1). Variations in corticosteroid sensitivity are also found in healthy people.

**Genetic factors**

Genetic variants of the glucocorticoid receptor (GR) gene are implicated in differences in corticosteroid sensitivity in healthy subjects and some inflammatory diseases, although GR polymorphisms have not been reported in corticosteroid resistant asthma. Single-nucleotide polymorphisms (SNPs) of genes affecting the corticosteroid pathway have been described in children with asthma recruited to the Childhood Asthma Management Program (CAMP) and in replication populations. The CAMP study followed children with asthma who were treated with inhaled corticosteroids over a 4-year period with efficacy outcomes including improvements in FEV₁, airway responsiveness to methacholine and the protection from exacerbations [19]. SNP variants associated with altered therapeutic response to treatment with inhaled corticosteroids include T-box expressed in T cells21 (TBX21) [20], corticotrophin-releasing hormone receptor 1 gene (CRHR1) [21], low-affinity IgE receptor gene (FCER2) [22]
and glucocorticoid-induced transcript 1 gene (GLCCI1) [23]. Genome-wide association studies (GWAS) have identified novel pharmacogenetic loci associated with variations in response to inhaled corticosteroids including T gene SNPs [24] and SNPs chr6 rs6924808 and chr11 rs1353649 [25]. Genes implicated in fetal lung development are also associated with corticosteroid treatment response [26]. The contribution of a single SNP to explaining corticosteroid insensitivity in the total asthma population is likely to be small, whereas combinations of polymorphisms in corticosteroid sensitive genes may have a more important effect on the therapeutic response to corticosteroids.

**Environmental factors**

**Cigarette smoke**

Corticosteroid insensitivity is an important clinical feature of current smokers with asthma [27-31]. Asthmatic smokers are less responsive to short and medium term treatment with inhaled or oral corticosteroids when assessed by improvements in symptoms, lung function and exacerbation rates compared to asthma patients who do not smoke [27-32]. Long-term treatment with inhaled corticosteroids however, may have beneficial effects in smokers with asthma by reducing the rate of decline in lung function [32-35]. Although insensitivity to inhaled corticosteroids is likely to contribute to poor asthma control in smokers, there are a number of unresolved issues regarding this phenomenon including the observation that not all smokers with asthma show insensitivity to short-term corticosteroid therapy, possibly due to differences in airway inflammation or intensity of smoking. The relationship between the intensity of exposure to cigarette smoke, either daily or cumulatively, and the development of corticosteroid insensitivity is not clearly established. Most studies that report a reduced
responsiveness to corticosteroids have recruited patients with a heavy smoking history of greater than 10 pack years.

A history of exposure to secondhand smoke in utero is associated with a reduced improvement in airway responsiveness to inhaled corticosteroids in children recruited to the CAMP study [36]. The mechanism underlying the attenuated response to inhaled corticosteroids in not known, although exposure to secondhand smoke in utero may adversely affect both the structure and function of the airways.

**Allergen exposure**

GR binding affinity of PBMCs from ragweed sensitive asthmatic subjects is reduced during the ragweed season when compared to before and after the pollen season [37]. In contrast, natural exposure to birch pollen does not reduce the ability of glucocorticoids to inhibit granulocyte-macrophage colony-stimulating factor production from PBMCs from subjects with seasonal allergic asthma and rhinitis due to birch pollen [38]. Severe asthma in children with fungal sensitization is associated with elevated airway interleukin (IL)-33 concentrations, which has been shown to induce corticosteroid insensitivity in a mouse model of asthma after exposure to the fungus *Alternaria alternata* [39]. Based on the current evidence, the importance of allergen exposure in corticosteroid insensitivity *in vivo* in asthma remains to be clearly established.

**Infections**

Toxins expression by microbes, called superantigens, are proposed as a possible mechanism by which infection could induce corticosteroid insensitivity [40]. Superantigens inhibit
corticosteroid-induced translocation of GRα to the cell nucleus [40] and increase the
numbers of nonfunctioning GRβ [41]. Alteration to the airway microbiome has been
implicated in corticosteroid responsiveness in asthma [42]. Corticosteroid-resistant asthma is
associated with classical antimicrobial activation of airway macrophages [43] and expansion
of specific gram-negative bacteria, which in vitro induce activate of mitogen-activated protein
kinase (MAP) and corticosteroid insensitivity [42]. In a mouse model, the microbial agonists
lipopolysaccharide (LPS) and β-glucan act synergistically with house dust mite to induce a
corticosteroid insensitive neutrophilic inflammation though toll-like (TL)-4 and dectin-1
receptors [44].

Virus-induced exacerbations including rhinovirus infection are insensitive to treatment with
corticosteroids, especially in children with asthma [45]. Rhinovirus infection causes
corticosteroid insensitivity in airway epithelium through induction of nuclear factor κB (NFκB)
and c-Jun N-terminal kinase (JNK) activation [46] and in monocyte cell lines through a
reduction in histone deacetylase (HDAC)2 activity [47]. The clinical importance of
superantigens, the airway microbiome and viruses in inducing corticosteroid resistance in
asthma remains to be established.

**Endotoxin**

Increased sputum endotoxin concentrations are associated with an impaired lung function
response to oral corticosteroids, particularly in never smokers with asthma, which suggests
that airway endotoxin may contribute to corticosteroid insensitivity in asthma [48] (Figure 2).

Patients with asthma may be exposed to endotoxin in household dust, through certain
occupations such as cotton textile workers or farm animal workers and in cigarette smoke,
either as active smokers or by exposure to secondhand smoke. Increased bacterial load within the airways or increased airway permeability might also increase airway endotoxin concentrations in asthma.

**Vitamin D deficiency**

Reduced vitamin D concentration in asthma are associated with reduced lung function, increased airway hyperresponsiveness and impaired corticosteroid sensitivity, when measured by dexamethasone-induced expression of mitogen-activated protein kinase phosphatase (MKP)1 by PBMCs [49].

**Asthma associated factors**

**Airway inflammatory cell phenotypes and airway remodeling**

Eosinophilic airway inflammation, identified by the presence of sputum or blood eosinophilia, predicts patients with asthma who are likely to obtain a favorable therapeutic response to corticosteroids. The type 2 helper T-cell (Th₂)-high subtype of asthma is associated with increased epithelial expression of interleukin (IL)-4, IL-5 and IL-13 [11] and predicts a beneficial therapeutic response to corticosteroids [11]. Many patients with asthma have non-eosinophilic asthma, sometimes associated with neutrophilic inflammation and/or have a Th₂-low type of inflammation. Non-eosinophilic inflammation is associated with an impaired therapeutic response to inhaled corticosteroids [50,51], although the lack of efficacy may not be complete. Intermittent eosinophilia might be a factor accounting for corticosteroid sensitivity in some of this group [51]. The numbers of peripheral blood CD8(+) T cells expressing the leukotriene B₄ (LTB₄) receptor BLT1 are increased in corticosteroid resistant
asthma [52]. Increased reticular basement membrane (RBM) thickness, which is a pathologic feature of airway remodelling, is associated with an impaired improvement in FEV$_1$ after a short course of systemic corticosteroids in severe asthma [53].

**Corticosteroid resistant asthma**

A very small proportion of patients with severe asthma have corticosteroid resistant asthma defined by less than a 15% improvement in FEV$_1$ after 2 weeks of prednisolone at a dose of 40 mg daily. Using an alternative definition of corticosteroid resistance, based on an increase in morning peak expiratory flow (PEF) of $\leq$ 15% following a course of oral corticosteroid, a retrospective study of 784 subjects with poorly controlled asthma found that 26% of subjects fulfilled this definition of corticosteroid resistant asthma [5].

**Non-adherence and poor inhaler technique**

Non-adherence with anti-asthma treatment is one of the most important reasons for poor symptom control and apparent corticosteroid insensitivity [54], with around a quarter of exacerbations estimated to be due to non-adherence with inhaled corticosteroids [55]. A high proportion of patients with difficult to control asthma have poor adherence with prescribed therapy, including both inhaled and oral corticosteroids [56]. Only a small percentage of inhaled corticosteroid reaches the target area in the lung, with the majority lost in the oropharynx or swallowed following inhalation. Good technique is paramount to ensure the benefits of inhaled corticosteroid treatment, although around half of all patients have poor inhaler technique irrespective of the device used [202].
**Concomitant disease and misdiagnosis**

Several concomitant conditions can contribute to poor symptom control or result in misdiagnosis and these should be considered in the event of a poor response to treatment including corticosteroids. For example, psychological factors are associated with worse clinical outcomes in asthma and psychosocial stress has been proposed to induce corticosteroid insensitivity in patients with asthma due to impaired GR expression and/or function [57]. The therapeutic response to corticosteroids in patients considered to have the asthma-COPD overlap syndrome is not clearly established, but in some patients there may be an impaired response.

**Demographic variables**

**Age and gender**

Age is positively correlated with increased in vitro peripheral blood T lymphocyte resistance to corticosteroids [58]. In Asthma Clinical Research Network (ACRN) trials in non-smokers with mild to moderate asthma, treatment failures (worsening asthma resulting in systemic corticosteroid use, hospitalization, emergency department visit, prolonged decrease in PEF, increase in albuterol use, or safety concerns including exacerbations) increased for every year above age 30 years in participants receiving inhaled corticosteroids [59]. The study did not specifically assess corticosteroid sensitivity. In the same analysis there was no difference in treatment failures between males and females [59].

**Race**
In black people with asthma the *in vitro* suppressive effects of dexamethasone on PBMC T lymphocyte proliferation and transcriptional gene expression responses are impaired compared to white people with asthma [16,58]. These findings suggest that a racial predisposition to impaired corticosteroid responsiveness could contribute to worse asthma control. In contrast, patients with mild to moderate asthma recruited to ACRN clinical trials treated with inhaled corticosteroids alone had no more treatment failures including exacerbations in African-Americans compared with white participants [60]. In a study of adolescents and adults with asthma, the proportion of African-Americans was not significantly associated with responsiveness to six weeks treatment with inhaled beclometasone [61].

**High BMI**

Mild or moderate asthmatic patients with a high body mass index (BMI) are less likely to achieve asthma control with inhaled fluticasone propionate treatment for 12 weeks compared to patients with a normal BMI [62] and have a smaller reduction in exhaled nitric oxide levels with inhaled corticosteroid treatment [63]. Dexamethasone induced MKP-1 expression *in vitro* is blunted in PBMCs and bronchoalveolar lavage (BAL) cells in overweight and obese patients with asthma [64]. A cluster analysis of adults with symptomatic airflow obstruction identified an obese-comorbid phenotype with late-onset asthma that obtained improvement in quality of life, although not in PEF variability, after inhaled corticosteroids [65].

**Pharmacokinetics of corticosteroids**
Impaired access of inhaled corticosteroids to target cells in the airways could result in corticosteroid insensitivity. Airway mucosal permeability, which is increased in both normal smokers and in asthma or excess mucus lining the airways of asthmatic smokers could impair the pharmacokinetics of inhaled corticosteroids. Dysfunction of small airways is thought contribute to poor symptom control in severe asthma and in smokers with asthma and due to the reduced deposition of large particle inhaled corticosteroids to the small airways might contribute to corticosteroid insensitivity. The particle size of dry powder fluticasone propionate is increased in the presence of tobacco smoke and based on this finding it has been postulated that cigarette smoking could prevent the dispersion of corticosteroid into the small airways of the lung and hence inhaler efficacy [66]. The pharmacokinetics of oral prednisolone in asthma is similar to healthy controls [67].

3. **Mechanisms of corticosteroids insensitivity**

Corticosteroids are the most effective treatment for asthma through their inhibitory actions on inflammatory cells and pathways involved in the pathogenesis of the disease [68]. Corticosteroids reduce inflammation through activation of GRs in the cytoplasm of target cells. The activated GR–corticosteroid complex binds to glucocorticoid-responsive elements (GREs) in the promoter region of corticosteroid-responsive genes within the nucleus to suppress (transrepression) or induce (transactivation) glucocorticoid target genes. Transrepression is believed to be responsible for the majority of the anti-inflammatory activity of corticosteroids through the reduced synthesis of proinflammatory transcription factors and by preventing the stimulation of inflammatory genes. The molecular mechanisms for the anti-inflammatory effects of corticosteroids are complex and not fully elucidated [69].
However, a key mechanism is thought to involve the reversal of histone acetylation of activated inflammatory genes through ligand-bound GRs binding to co-activator molecules and subsequent recruitment of HDAC2 to the activated inflammatory gene transcription complex [68]. Recent studies suggest that the production of anti-inflammatory proteins may have a more important role in resolution of inflammation than previously considered [69]. Corticosteroids induce histone acetylation of anti-inflammatory genes (transactivation), including MKP-1 and glucocorticoid-inducible leucine zipper 1 (GILZ-1) as well as potentiating the effects of \( \beta_2 \) agonists, in part through increased expression of \( \beta_2 \) adrenergic receptors on cell surfaces [68]. The adverse effects of corticosteroids are thought to be due to transactivation of genes involved in glucose, lipid and muscle metabolism as well as by interacting with negative GREs to inhibit the production of osteocalcin, which is involved in bone synthesis [68]. Corticosteroids also exert anti-inflammatory effects through a non-genomic mechanism, which has a rapid onset within minutes, is of short duration and is dose-dependent. Non-genomic mechanisms are thought to include activation of endothelial nitric oxide synthase (eNOS) resulting in the production of nitric oxide (NO) and the release of the anti-inflammatory molecule annexin-1.

Several mechanisms could explain the variation between people in the response to corticosteroids including differences in genetic, inflammatory cell phenotypes and other factors discussed in section 2. Molecular mechanisms of corticosteroid insensitivity in asthma have been extensively reviewed in previous publications [70-72] and only a summary of the key mechanisms are reviewed in this article. Postulated molecular mechanisms for corticosteroid insensitivity in asthma include alteration in GR subtypes, defective GR binding
and impaired GR nuclear translocation, increased proinflammatory transcription factors and defective histone acetylation (Table 2).

**Alteration in glucocorticoid receptor subtypes**

GRα is a functional receptor that binds corticosteroids to induce or suppress the transcription of multiple genes and is the predominant GR in most inflammatory cells. Isoforms of GR are derived by alternative splicing and one subtype GRβ is unable to bind corticosteroids. It has been proposed that corticosteroid insensitivity in asthma occurs due to over expression of the non-functional GRβ subtype or a reduction in ligand-activated GRα subtype numbers or activity. GRβ expression on cells is increased by several factors associated with corticosteroid insensitivity in asthma including superantigens [41] and proinflammatory mediators [73] as well as by cigarette smoking, where the ratio of GRα to GRβ in PBMCs is reduced in cigarette smokers compared with never smokers [74]. Elevated GRβ expression in corticosteroid-resistant asthma may reduce nuclear translocation of GRα and decrease HDAC2 expression in response to corticosteroids [75]. Overall the data is conflicting on the importance of alterations in the GRβ subtype in causing corticosteroid insensitivity in asthma [75,76]. A cluster analysis of patient with mild to moderate asthma reported that obese subjects have reduced expression of GCRα in PBMCs that correlates with a reduced induction of MKP-1 expression by dexamethasone [77].

**Defective glucocorticoid receptor binding and nuclear translocation**
A reduction in GR binding affinity and nuclear translocation could contribute to corticosteroid insensitivity through the action of pro-inflammatory mediators, such as IL-2 and IL-4 in combination or IL-13. Cytokines are thought to induce these effects by phosphorylation of GR through the activation of protein kinases including p38 MAPK, JNK and extracellular signal-regulated kinase (ERK) [71]. Corticosteroids activate endogenous inhibitors of p38MAPK and JNK pathways including MKP-1 and protein phosphatase 2A (PP2A) [71,78]. The expression of the anti-inflammatory protein MKP-1 is reduced in alveolar macrophages from patient with severe asthma associated with corticosteroid insensitivity [79] and in PBMCs from overweight and obese patients with asthma [64,77]. In a mouse model, the translocation of GR to the nucleus is inhibited by IFN-γ producing cells and LPS cooperating to produce IL-27 from macrophages through activation of toll-like receptor (TLR)4 and MyD88 dependent pathways [80]. Furthermore, IFN-γ and LPS induce miR-9 expression in lung tissue and primary macrophages, which reduces PP2A activity [81]. Corticosteroid induced GRα nuclear translocation is suppressed by rhinovirus infection in airway epithelium through JNK activation [46]. Recently asthmatic children clinically classified as poor corticosteroid responders were reported to have reduced GR bioavailability due to decreased GR protein expression and more rapid corticosteroid-induced down regulation compared to asthmatic children classified as good corticosteroid responders [82].

**Increase in proinflammatory transcription factors**

Activation of pro-inflammatory transcription factors such as NF-κB, activator protein-1 (AP-1) and signal transduction-activated transcription factors (STAT) enable the translation of inflammatory genes and recruiting of inflammatory cells and could cause corticosteroid
insensitivity. Increased nuclear factor-κB (NF-κB) expression in PBMCs has been associated with corticosteroid insensitivity in severe asthma [83] and as tobacco smoke activates NF-κB, this transcription factor which might contribute to reduced response to corticosteroids in smokers with asthma. Dysregulation of AP-1 is associated with corticosteroid resistant asthma [84]. Rhinovirus infection induce corticosteroid resistance in airway epithelium by the activation of NF-κB [46].

**Defective histone acetylation**

Corticosteroids require HDAC activity for maximal suppression of inflammatory cytokine induction [85]. Low levels of HDAC2 expression are found in PBMCs [86] and alveolar macrophages [87] in patients with severe asthma, which is thought to be due to excessive oxidative stress [71]. Smokers have decreased HDAC2 activity in alveolar macrophages, possibly as a result of oxidative and nitrative stress and this may lead to increased inflammatory gene expression and reduced sensitivity to corticosteroids [88,89]. Oxidative stress activates phosphinositide 3-kinase (PI3K) resulting in AKt phosphorylation and HDAC2 inactivation [71].

**Other immune mechanisms**

Suppression of the anti-inflammatory cytokine IL-10 production by T lymphocytes in asthma could contribute to corticosteroid insensitivity [90]. The pro-inflammatory mediator TNFα induces corticosteroid insensitivity in a mouse model of neutrophilic airway inflammation [91].
4. Current management of corticosteroid insensitivity

The aims of management are to achieve good asthma control and to minimize treatment burden, especially from oral corticosteroid use, although achieving these aims in severe asthma is often difficult. A step-wise approach to treatment should be integrated into a management plan that includes assessment of adherence and inhaler technique, non-pharmacological treatments where appropriate, as well as supported self-management [201, 202]. It is important to distinguish patients with severe asthma from those with ‘difficult to control’ asthma due to reasons other than disease severity by undertaking a systematic evaluation [92]. For adults and adolescents with asthma patients who have poor symptom control or who at risk of exacerbations and in whom the diagnosis of asthma has been confirmed, guidelines recommend daily inhaled corticosteroid as the preferred controller treatment, with the addition of add-on therapies if symptoms remains poorly controlled [201, 202]. However, this approach may result in up to one third of patients failing to obtain maximal improvement in asthma control and quality of life due to corticosteroid insensitivity. Several management strategies that address specific factors associated with an impaired therapeutic response to corticosteroids may result in improvements in asthma control (Table 3).

Non-adherence

The ability to detect poor adherence based on history alone is poor. Adherence can be assessed by reviewing prescription refill frequency and by measuring circulating blood prednisolone concentration in patients taking oral corticosteroids. A trial of a parenteral depo corticosteroid can help establish whether persistent symptoms are due to poor adherence with inhaled and/or
oral corticosteroids. Strategies to improve adherence include adequate explanation of
the indications for treatment, discussion of real and perceived concerns of adverse effects of
treatment, simplifying drug treatment regimens, reminders and reinforcement [54]. Improving
medication adherence could have a substantial beneficial impact on asthma related clinical
outcomes [93].

**Non-pharmacological interventions**

There is limited information on the effectiveness of non-pharmacological interventions for the
management of factors associated with corticosteroid insensitivity in asthma (Table 1). A small
number of studies have examined the role of smoking cessation on asthma outcomes and
reported improvements in symptoms and lung function in those people who quit smoking
successfully [30]. Smoking cessation in asthma is associated with improvement in corticosteroid
sensitivity [15,94]. Weight reduction in asthmatic patients with a high BMI may result in
improvements in asthma symptoms [95], although whether corticosteroid sensitivity also
improves is not reported.

Anti-inflammatory and corticosteroid-enhancing actions of vitamin D are reported in monocytes of
patients with corticosteroid-resistant asthma and those with steroid-sensitive asthma, although
the responses to corticosteroids remains lower than in patients with corticosteroid sensitive
asthma [96,97]. Peripheral blood CD4+ T cells isolated from subjects with corticosteroid-resistant
asthma regain their sensitivity to corticosteroids following a short course of oral vitamin D through
the induction of IL-10 [98]. Based on these findings, several clinical trial have examined the effect
of dietary supplement of vitamin D in asthma. A small clinical trial of vitamin D3 supplementation
in patients with corticosteroid resistant asthma demonstrated a modest improvement in lung function after a short course of oral prednisolone and an increase in dexamethasone-induced IL-10 production \textit{in vitro} as well as suppression of dexamethasone-induced IL-17A production [99,100]. Two large randomized clinical trials of vitamin D3 supplementation in patients with asthma and vitamin D insufficiency [VIDA and ViDiAs trials), although not selected for corticosteroid insensitivity, reported no improvements in clinical outcome [101,102]. Interestingly, vitamin D supplementation reduced eosinophilic inflammation in patients with non-atopic asthma, suggesting that certain phenotypes might obtain benefit from vitamin D3 supplementation [103].

**Current pharmacological treatments**

A step-up in the dose of corticosteroid as well as the addition of other therapies may benefit individuals with asthma that is associated with corticosteroid insensitivity.

**Inhaled corticosteroids**

The relatively flat dose response to inhaled corticosteroids in asthma results in the majority of the therapeutic benefits being achieved at low to medium doses [1]. In the Gaining Optimal Asthma Control (GOAL) trial, high dose inhaled corticosteroids treatment with inhaled fluticasone propionate 1000 mcg daily resulted in clinical improvements in asthma control, although in patients with more severe disease only 50% achieved well controlled asthma and 20% achieve totally control [2] [201]. In a short-term study in smokers with mild asthma who demonstrated corticosteroid insensitivity to low dose inhaled corticosteroids there was an improvement in lung function with high dose inhaled corticosteroid treatment [31]. Targeting the small airways with extrafine-particle inhaled corticosteroids may
potentially impact favorably on the safety and efficacy of inhaled corticosteroids through improved total lung deposition and resulting in improved asthma control at lower daily doses than the large-particle inhaled corticosteroid. Observational studies suggest that extrafine particle HFA-BDP Qvar® may achieve better control at lower prescribed doses than large particle inhaled corticosteroids [104], but to date randomized controlled trials have not clearly demonstrated that equipotent doses of small-particle inhaled corticosteroids are more effective than large-particle inhaled corticosteroids.

**LABAs in combination with inhaled corticosteroids**

The combination of a LABA and an inhaled corticosteroid may act synergistically to increase corticosteroid sensitivity of inflammatory cell in severe asthma and in smokers with asthma. Long acting β₂-agonists have been shown *in vitro* to increase the translocation of GRs to the nucleus. The LABA formoterol increases corticosteroid sensitivity *in vitro* through inhibition of GR phosphorylation via PP2A activation in a β₂-receptor independent manner [105] and by inhibition of PI3K signaling [106]. In addition, formoterol enhances the expression of the anti-inflammatory transcription factor MKP-1 induced by corticosteroids [107].

**Low dose theophylline**

Low dose theophylline restores corticosteroid sensitivity *in-vitro* possibly by increasing HDAC2 activity, which is suppressed in severe asthma and in smokers and a similar clinical effect might occur in asthma[108]. An exploratory study examined the impact of low dose theophylline added to inhaled corticosteroid compared to inhaled corticosteroid alone in a group of smokers with asthma [109] (Figure 3). The addition of theophylline to inhaled corticosteroid, at a dose titrated to provide a ‘sub-therapeutic’ concentration, resulted in
increased efficacy as measured by lung function and suggested the restoration of corticosteroid sensitivity in those treated with the combination. Theophylline inhibits oxidative stress dependent PI3K-δ activation and restores corticosteroid sensitivity in PBMCs from patients with COPD [108]. Clinical trials have not investigated the therapeutic effects of adding low dose theophylline in patients with severe asthma. A fixed combination of ultra-low dose of theophylline with fluticasone, SKP-2075 is under development for the treatment of COPD. This combination would potentially be of benefit in the treatment of corticosteroid insensitivity in severe asthma and in smokers with asthma.

**Leukotriene-receptor antagonist**

A randomized controlled trial that compared oral montelukast (10 mg daily), inhaled fluticasone propionate (500 µg daily), and placebo over a 6-month treatment period in mild smokers with asthma [110] reported that less than 50 percentage of days with asthma control during each treatment, although both active treatments were better than placebo. Patients with a smoking history of ≤11 pack years tended to show greater benefit with fluticasone, whereas those with a smoking history of >11 pack years tended to show greater benefit with montelukast.

**Oral corticosteroid sparing therapies**

Some biological agents and immunosuppressive agents have oral corticosteroid sparing effects in patients with severe asthma who have uncontrolled symptoms despite treatment that includes high dose inhaled corticosteroids and LABAs, although it is unclear whether the reduction in daily oral corticosteroids is due to a reversal of corticosteroid insensitivity or due to the intervention targeting corticosteroid insensitive inflammatory pathways. Omalizumab is a humanised
monoclonal antibody that binds circulating IgE antibody produces improvements in quality of life and reductions in exacerbations [111] as well as possibly having oral corticosteroid-sparing effects. An anti-IL-5 monoclonal antibody mepolizumab, which blocks the binding of IL-5 to IL-5 receptor-α on the surface of eosinophils, decreases severe exacerbations [112] and reduces the use of oral corticosteroids [113]. Mepolizumab was recently licensed in the US for use in patients with severe eosinophilic asthma.

5. Potential future therapies for corticosteroid insensitivity

Many patients with asthma that is associated with corticosteroid insensitivity continue to have poorly controlled disease despite treatment with currently available therapies. There is an unmet need for new treatments that will impact favorably on clinical outcomes in these patients through the restoration of corticosteroid sensitivity (Table 4). Novel small molecule drugs and biologics agents have been developed as possible new treatments for asthma, including for patients with corticosteroid insensitivity. Several drugs licensed for the treatment of medical conditions other than asthma have been investigated for their efficacy in asthma. Candidate drugs have been chosen usually because of pre-clinical evidence of effects on reduction inflammation and restoring corticosteroid sensitivity that might be relevant to the treatment of asthma. Potential therapies for patients with severe refractory asthma associated with corticosteroid insensitivity are summarized below.
Small molecule drugs

**Macrolides**

The mechanism(s) of action of macrolides in the treatment of airway diseases is not established, but could be due to their antibacterial and/or anti-inflammatory actions, which include inhibition of NF-κB and other transcription factors as well as reduction in IL-8 production, neutrophil migration and/or function [114,115]. Additionally, macrolides may restore corticosteroid sensitivity by inhibiting PI3K activation [116] and TNFα/IL-17 immune responses [117]. Two recent exploratory clinical trials have investigated the effects of macrolides in non-eosinophilic asthma associated with corticosteroid insensitivity. In the first trial, seventy-seven smokers with mild to moderate asthma were randomized to a double-blind parallel-group trial comparing azithromycin [250 mg per day] with placebo [118]. At 12 weeks, treatment with azithromycin did not improve symptom control or lung function. In the second randomized controlled trial non-smokers with exacerbation-prone severe asthma received low-dose azithromycin (n=55) or placebo (n=54) as add-on treatment to combination therapy of inhaled corticosteroids and LABA for 6 months [119]. Azithromycin did not reduce the rate of severe exacerbations and lower respiratory tract infections requiring treatment with antibiotics in the total study population. In a predefined subgroup there was a reduction in the rate of primary endpoints in azithromycin-treated patients with non-eosinophilic severe asthma (blood eosinophilia ≤200/µl) [119]. Novel analogue of macrolides have been developed that have greater anti-inflammatory effects than current macrolides, such as solithromycin (CEM-101) [115,116] or that lack anti-bacterial properties but retain anti-inflammatory activity, such as the non-antibiotic azithromycin derivative CSY0073 [120].
**Protein kinase inhibitors**

Inhibition of protein kinase, such as p38MAPK, narrow spectrum kinase and tyrosine kinase, that are involved in cellular signaling of pro-inflammatory cytokines, may have a role in the treatment of severe asthma associated with corticosteroid insensitivity [121-123]. Several p38MAPK inhibitors restore corticosteroid sensitivity in PBMCs from patients with severe asthma [121,124] and COPD [125]. Clinical trials of p38MAPK inhibitors, such as oral losmapimod (GW856553) and inhaled AZD7624 are underway in COPD, although none are registered in asthma. A specific ckit tyrosine kinase inhibitor imatinib inhibits hyperresponsiveness, inflammation and remodeling in murine asthma models [126] and is undergoing clinical trial evaluation in severe refractory asthma (ClinicalTrials.gov Identifier: NCT01097694). A tyrosine kinase inhibitor masitinib that targets c-kit and the platelet-derived growth factor (PDGF) receptor improved asthma control in patients with severe corticosteroid-dependent asthma [127] and a further clinical trial is underway in patients with severe persistent asthma treated with oral corticosteroids (ClinicalTrials.gov Identifier: NCT01449162).

**PI3kinase inhibitors**

Pre-clinical studies suggest that PI3K-δ inhibitors could potentially reverse corticosteroid insensitivity by increasing HDAC2 activity [128,129]. Selective PI3K inhibitors are being developed as novel treatments of corticosteroid insensitive airway diseases. An inhaled PI3Kδ inhibitor GSK2269557 is undergoing clinical trials in adults with persistent uncontrolled asthma (ClinicalTrials.gov Identifier: NCT02567708). PI3K δ and γ isoforms are implicated in inflammatory cell recruitment and activation [129]. An inhaled dual PI3Kδ/γ inhibitor, TG100-115, reduces airway inflammation induced by allergen and cigarette smoke in murine models.
and restores corticosteroid sensitivity in the smoke model [130]. A PI3Kδ/γ inhibitor RV1729 has undergone early stage clinical evaluation in COPD, although no results are published (ClinicalTrials.gov Identifier: NCT02140346). The tricyclic antidepressant nortriptyline reverses corticosteroid insensitivity induced by oxidative stress in vitro by inhibition of PI3K-δ and it has been proposed as a potential treatment for corticosteroid-insensitivity in severe asthma, smokers with asthma and COPD [131].

**NF-κB inhibitors**

Targeting activation of the pro-inflammatory transcription factor NF-κB with inhibitors of IκB kinases (IKKs) may restore corticosteroid sensitivity in severe asthma and smokers with asthma [132], although adverse effects may limit the use of these compounds.

**PDE4 inhibitors and dual PDE3 and PDE4 inhibitors**

Phosphodiesterase (PDE)4 inhibitors have immunomodulatory properties relevant to the treatment of asthma including the restoration of corticosteroid sensitivity [133]. The oral PDE4 inhibitor roflumilast reverses corticosteroid resistance in peripheral blood neutrophils from patients with COPD [134]. High doses of PDE4 inhibitors may be necessary to treat severe asthma, although gastro-intestinal side effects limit their use. Inhaled formulations may improve the therapeutic ratio of PDE4 inhibitors [135-137]. The inhaled PDE4 inhibitor GSK256066 inhibits allergen-induced late asthmatic responses [136] and in patients with moderate COPD GSK256066 was well tolerated although there was no inhibitory effect on inflammatory biomarkers in sputum and blood [138]. The combination of selective PDE3 and PDE4 inhibitors prevent corticosteroid insensitivity induced by oxidative stress in human alveolar macrophages [139].
inhaled dual PDE\textsubscript{3} and PDE\textsubscript{4} inhibitor, RPL554 has bronchodilator effects and is well tolerated in patients with asthma and COPD [140].

**PPAR\textgamma\, agonists**

Peroxisome proliferator-activated receptor-\textgamma\,(PPAR\textgamma) agonists are members of the nuclear receptor family that exert in pre-clinical studies anti-inflammatory effects potentially relevant to the treatment of airway diseases and to restoring corticosteroid sensitivity [141,142]. A proof of concept study using the oral PPAR\textgamma agonist rosiglitazone demonstrated bronchodilator effects in mild to moderate smokers with asthma who had non-eosinophilic inflammation [143]. A borderline reduction in sputum IL-8 was observed with rosiglitazone compared with inhaled beclometasone dipropionate. The adverse-effect profile of oral rosiglitazone has precluded its development as a treatment for asthma. Inhaled PPAR\textgamma agonist analogues, such as ADC3277 should facilitate further examination of this potential novel approach to the treatment of corticosteroid insensitive airway disease.

**Statins**

Statins reduce cholesterol biosynthesis by inhibiting 3-hydroxy-3-methylglutaryl coenzyme A (HMG-CoA) reductase and in addition have pleiotropic immunomodulatory effects that may be of value for the treatment of chronic inflammatory diseases [144,145]. Preclinical in vitro and in vivo studies, including experimental models of allergic [146] and tobacco-smoke-induced lung inflammation [147] found that statins reduce components of airway inflammation potentially relevant to the pathogenesis of asthma and smoke-induced airway diseases. Statins may also restore corticosteroid sensitivity in asthma [148,149]. In smokers with asthma the effects of atorvastatin alone and in combination with inhaled corticosteroid
was investigated on the ability to suppress the concentration of a range of cytokines, chemokines and growth factors in sputum [150,151]. Despite the absence of a suppressive effect of inhaled beclometasone on sputum mediators, atorvastatin either alone or in combination with inhaled beclometasone reduced sputum concentrations of approximately one-third of the selected cytokines, chemokines and growth factors. Atorvastatin significantly reduced sputum concentrations of CCL7, IL-12p70, sCD40L, FGF-2, CCL4, TGF-α and MMP-8 compared with placebo and, when combined with inhaled beclometasone, reduced sputum concentrations of MMP-8, IL-1β, IL-10, MMP-9, sCD40L, FGF-2, IL-7, G-CSF and CCL7 compared to inhaled beclometasone alone (Figure 4). Improvements in ACQ and/or AQLQ scores with atorvastatin and inhaled corticosteroid were associated with decreases in G-CSF, IL-7, CCL2 and CXCL8. Interestingly, simvastatin suppresses airway IL-17 and upregulated IL-10 in patients with stable COPD [152]. Taken together, these findings suggest that treatment with atorvastatin either alone or in combination with an inhaled corticosteroid may be of benefit in certain corticosteroid insensitive airway diseases. There is a need for long-term clinical studies examining effect of statins on exacerbations, neutrophil function and airway remodeling in severe asthma associated with corticosteroid insensitivity.

**Drugs targeting neutrophilic inflammation**

As increased airway neutrophils have implicated as a potential cause corticosteroid insensitivity in asthma, small molecule drugs have been developed to treat asthma that target these cells including CXCR2 antagonists in asthma [153,154] and COPD [155] and a 5-lipoxygenase-activating protein (FLAP) inhibitor GSK2190915 in asthma [156], although their ability to restore corticosteroid sensitivity has not been studied. In addition, evidence for the
clinical effectiveness of CXCR2 antagonists and FLAP inhibitors in chronic asthma is so far lacking.

**Biological agents**

Pre-clinical studies suggest a possible role for Th\textsubscript{17} cells and IL-17 in neutrophilic asthma and corticosteroid insensitivity [157]. Increased levels of IL-17 are found in BAL, sputum and bronchial biopsy samples obtained from patients with severe asthma and correlates with sputum neutrophils. Brodalumab is a human monoclonal antibody that binds with high affinity to human IL-17RA, blocking the biologic activity of IL-17A, -17F, -17A/F heterodimer, and IL-25. A randomized clinical trial of brodalumab administered for 12 weeks to 302 adults with inadequately controlled moderate to severe asthma taking regular inhaled corticosteroids found no treatment difference in ACQ score or in secondary endpoints including FEV\textsubscript{1} and symptom-free days [158]. In a pre-specified subgroup with high bronchodilator reversibility that accounted for 37% of the full study population there was a borderline improvement an ACQ score. The clinical trial did not select patients with neutrophilic inflammation. A follow-up clinical trial of brodalumab in inadequately controlled asthma subjects with high bronchodilator reversibility was recently terminated (ClinicalTrials.gov Identifier: NCT01902290). A clinical trial of the IL-17A monoclonal antibody blocker secukinumab in patients with uncontrolled asthma has also been recently terminated (ClinicalTrials.gov Identifier: NCT01478360). Neutralizing TNF\textsubscript{α} restores corticosteroid sensitivity in a mouse model of neutrophilic airway inflammation [91]. Despite some promise shown in early small clinical studies with the soluble TNF-\(\alpha\) receptor blocker etanercept in severe asthma, larger studies with golimumab [159] and etanercept [160] did not confirmed a
consistent effect. Overall, when combined with concerns over increased risk of severe
infections and malignancies with treatment [159] it is unlikely that TNF-α receptor blockers
will be developed further for the treatment of asthma. Monoclonal antibodies that block
IL-1β, for example, canakinumab might be of benefit in corticosteroid insensitive asthma,
although no clinical studies are currently registered.

6. Expert commentary

Corticosteroids are the most effective treatment for asthma, but the therapeutic response
differs markedly between individuals. Corticosteroid insensitivity may be clinically important
in up to a third of patients with asthma particularly in people with severe disease. Multiple
factors including genetic, environmental such as cigarette smoking and asthma-related such
as non-adherence and non-eosinophilic inflammation as well as demographic variables are
likely to account for the heterogeneous response to corticosteroids between people.

Advances in the understanding the molecular mechanisms of corticosteroids have led to
important insights into the pathways involved in corticosteroid insensitivity in asthma
including alterations in GR subtypes, defective GR binding and nuclear translocation,
increased proinflammatory transcription factors and defective histone acetylation as well as
other immunological mechanisms. Nevertheless, the underlying mechanisms of corticosteroid
insensitivity in asthma are only partly understood. In the future system biology approaches
may produce new insights to understanding corticosteroid insensitivity [72].

Non-adherence with inhaled and oral corticosteroid treatment is one of the most important
reasons for poor symptom control and apparent corticosteroid insensitivity. There is limited
information on the effectiveness of non-pharmacological interventions for the management of asthma on specific factors associated with corticosteroid insensitivity in asthma. Advice on smoking cessation is essential in smokers with asthma and may improve corticosteroid responsiveness.

A step-up in the dose of corticosteroid as well as the addition of other therapies for asthma may benefit some individuals with asthma that is associated with corticosteroid insensitivity. High dose inhaled corticosteroids treatment improves clinical outcomes in some patients with severe asthma and in smokers with asthma, but many patients remain symptomatic. Long acting $\beta_2$-agonists have been shown in vitro to increase the translocation of GR to the nucleus and to be of clinical benefit. In an exploratory clinical trial in smokers with asthma low dose theophylline added too inhaled corticosteroid resulted in improvement in lung function and suggested the restoration of corticosteroid sensitivity in those treated with the combination. It is not known whether a similar beneficial effect is achieved with low dose theophylline in severe asthma.

Some biological agents such as omalizumab and mepolizumab and immunosuppressive agents such as methotrexate have oral corticosteroid sparing effects in patients with severe asthma although it is not known whether the reduction in daily oral corticosteroids is due to a reversal of corticosteroid insensitivity.

Many patients with asthma continue to have poorly controlled disease that is associated with corticosteroid insensitivity despite receiving treatment with currently available therapies. Small molecule drugs such as macrolides, various protein kinase inhibitors, PI3 kinase inhibitors, NF-κB inhibitors, PDE$_4$ and PDE$_3$/PDE$_4$ inhibitors, PPAR$\gamma$ agonists as well as statins might be shown to be of benefit in the treatment of corticosteroid insensitive airway diseases. The results of clinical
trials of biological agents such as monoclonal inhibitors of IL-17 has been disappointing to date. Future clinical trials will determine which of these therapies will ultimately be licensed for the treatment of asthma and in particular will be of value in the treatment of corticosteroid insensitivity.

7. Five-year review

The management of patients with asthma is often hampered because it is currently difficult to identify whether poorly controlled symptoms are due to corticosteroid insensitivity or to other reasons such as non-adherence or the involvement of corticosteroid insensitive inflammatory pathways. The development of biomarkers that can identify corticosteroid insensitivity in the clinic would greatly aid in the management of patients with uncontrolled asthma and lead to a personalized approach to treatment. Over the last 15 years several international collaborative programmes of research have investigated pathogenic mechanism of severe asthma and corticosteroid insensitivity [161,162]. A recent international collaborative project in severe asthma, the Unbiased Biomarkers for the Prediction of Respiratory Disease Outcome (U-BOPRED) study, is designed to identify new phenotypes/endotypes and treatment targets using omics technologies and applying a system biology approach. It is due to report its main findings shortly [163]. The UK Refractory Asthma Stratification Programme (RASP-UK) is investigating novel biomarker stratification strategies in severe asthma, assessing adherence to corticosteroids and examining biomarkers to optimise corticosteroid treatment [164]. Hopefully the findings from these research networks will identify new approaches to the treatment of patients with severe corticosteroid insensitive asthma.
Key issues

• Corticosteroids are the most effective treatment for asthma, but the therapeutic response varies markedly between individuals, with up to one third of patients showing evidence of insensitivity to corticosteroids.

• Multiple factors including genetic, environmental such as cigarette smoking and asthma-related factors such as non-adherence and non-eosinophilic inflammation as well as demographic and pharmacokinetic variables are likely to contribute to the heterogeneous response to corticosteroids between people.

• Molecular mechanisms proposed to explain corticosteroid insensitivity include alterations in GR α and β subtypes, impaired GR binding and nuclear translocation, increased proinflammatory transcription factors and defective histone acetylation.

• Non-adherence with inhaled and oral corticosteroid treatment is one of the most important reasons for poor symptom control and apparent corticosteroid insensitivity.

• There is limited information on the effectiveness of non-pharmacological interventions for the management of asthma on specific factors associated with corticosteroid insensitivity in asthma. Advice on smoking cessation is essential in smokers with asthma and may improve corticosteroid responsiveness.

• A step-up in the dose of corticosteroid as well as the addition of other therapies for asthma may benefit some individuals with asthma that is associated with corticosteroid insensitivity. High dose inhaled corticosteroids treatment improves clinical outcomes in some patients with severe asthma and in smokers with asthma, but many patients remain
symptomatic. Long acting $\beta_2$-agonists have been shown \textit{in vitro} to increase the translocation of GR to the nucleus. In an exploratory clinical trial in smokers with asthma, low dose theophylline added to inhaled corticosteroid resulted in improvement in lung function and suggested the restoration of corticosteroid sensitivity in those treated with the combination.

- Future interventions that may restore corticosteroid sensitivity in asthma include small molecule drugs such as macrolides, several protein kinase inhibitors, PI3 kinase inhibitors, NF-$\kappa$B inhibitors, PDE$_4$ and PDE$_3$/PDE$_4$ inhibitors, PPAR$_\gamma$ agonists as well as statins and biological agents.

- In the future, biomarkers may be used in the clinic to predict corticosteroid insensitivity in patients with poorly controlled asthma.
KEY REFERENCES

Papers of special note have been highlighted as:

• of interest

•• of considerable interest


** A composite phenotype that includes information on inhaled corticosteroid treatment response in patients with mild to moderate asthma assessed by six outcomes (lung function, bronchodilator response, airway responsiveness, symptoms, need for oral corticosteroids and frequency of emergency department visits and hospitalizations) is reported to identify endophenotypes that more accurately describe corticosteroid sensitivity and insensitivity.


** Provides an overview of recent evidence on the harmful effects of smoking in asthma, possible underlying inflammatory mechanisms for this altered response, management options for these patients and potential future therapeutic directions.


46. Papi A, Contoli M, Adcock IM et al. Rhinovirus infection causes steroid resistance in
airway epithelium through nuclear factor κB and c-Jun N-terminal kinase activation. J

47. Footitt J, Mallia P, Durham AL et al. Oxidative and nitrosative stress and histone
deacetylase-2 activity in exacerbations of chronic obstructive pulmonary disease.

sputum endotoxin levels are associated with an impaired lung function response to
* Increased sputum endotoxin concentrations are associated with an impaired lung
function response to oral corticosteroids, particularly in never smokers with asthma,
which suggests that airway endotoxin may contribute to corticosteroid insensitivity in
asthma

49. Sutherland ER, Goleva E, Jackson LP, Stevens AD, Leung DYM. Vitamin D Levels, Lung

50. Pavord I, Brightling C, Woltmann G, Wardlaw A. Non-eosinophilic corticosteroid

51. McGrath KW, Icitovic N, Boushey HA et al. A Large Subgroup of Mild-to-Moderate
Asthma Is Persistently Noneosinophilic. Am J Respir Crit Care Med, 185(6), 612-619
(2012).

52. Chung EH, Jia Y, Ohnishi H et al. LeukotrieneB4 receptor 1 is Differentially Expressed
on Peripheral T Cells of Steroid-Sensitive and -Resistant Asthmatics. Ann Allergy

*Chest*, 141(6), 1504-1511 (2012).

54. Heaney LG, Horne R. Non-adherence in difficult asthma: time to take it seriously. 


* In Asthma Clinical Research Network (ACRN) trials in non-smokers with mild to moderate asthma, treatment failures (worsening asthma resulting in systemic corticosteroid use, hospitalization, emergency department visit, prolonged decrease in PEF, increase in albuterol use, or safety concerns including exacerbations) increased for every year above age 30 years in participants receiving inhaled corticosteroids.


* Comprehensive review of the mode of action of corticosteroids in asthma and COPD


** Comprehensive review of the mechanisms of corticosteroid resistance in asthma and COPD **


* Vitamin D3 supplementation in patients with corticosteroid resistant asthma demonstrated an increase in dexamethasone-induced IL-10 production in vitro as well as suppression of dexamethasone-induced IL-17A production


* An exploratory study examined the impact of low dose theophylline added to inhaled corticosteroid compared to inhaled corticosteroid alone in a group of smokers with asthma. The addition of theophylline to inhaled corticosteroid, at a dose titrated to provide a ‘sub-therapeutic’ concentration, resulted in increased efficacy as measured by lung function and suggested the restoration of corticosteroid sensitivity in those treated with the combination.


123. Chung KF. p38 Mitogen-Activated Protein Kinase Pathways in Asthma and COPD. *Chest*, 139(6), 1470-1479 (2011).


URL: https://mc.manuscriptcentral.com/errx   Email: Joseph.Walsh@informa.com

* Comprehensive review of selective PDE inhibitors as novel treatments for respiratory diseases.


140. Franciosi LG, Diamant Z, Banner KH et al. Efficacy and safety of RPL554, a dual PDE3 and PDE4 inhibitor, in healthy volunteers and in patients with asthma or chronic...

141. Spears M, McSharry C, Thomson NC. Peroxisome proliferator-activated receptor-

gamma agonists as potential anti-inflammatory agents in asthma and chronic


receptor-γ ligands on in vitro and in vivo models of COPD. *Eur Respir J*, 43(2), 409-420

(2014).


144. Hothersall E, McSharry C, Thomson NC. Potential therapeutic role for statins in


145. Yeganeh B, Wiechec E, Ande S et al. Targeting the mevalonate cascade as a new

therapeutic approach in heart disease, cancer and pulmonary disease. *Pharmacol

Ther*, 143(1), 87-110 (2014).

146. McKay A, Leung BP, McInnes IB, Thomson NC, Liew FY. A Novel Anti-Inflammatory

Role of Simvastatin in a Murine Model of Allergic Asthma. *J Immunol*, 172(5), 2903-

2908 (2004).


and chemokine production by peripheral blood mononuclear cells in patients with


* Short-term treatment with atorvastatin alone or in combination with inhaled beclometasone reduces several sputum cytokines, chemokines and growth factors concentrations unresponsive to inhaled corticosteroids alone, in smokers with asthma.


Websites

201 GINA Report, Global Strategy for Asthma Management and Prevention 2014


Table 1: Factors that potentially influence the therapeutic response to corticosteroids in patients with asthma

Genetic factors
- T-box expressed in T cells (TBX21) polymorphisms
- Corticotrophin-releasing hormone receptor 1 gene (CRHR1) polymorphisms
- Low-affinity IgE receptor gene (FCER2) polymorphisms
- Glucocorticoid-induced transcript 1 gene (GLCCI1) polymorphisms
- T gene polymorphisms
- chr6 rs6924808 and chr11 rs1353649 polymorphisms

Environmental factors
- Active and secondhand exposure to cigarette smoke
- Allergen exposure
- Infections: superantigens, airway microbiome, viruses
- Endotoxin
- Vitamin D deficiency

Asthma associated factors
- Airway inflammatory cell phenotypes and airway remodeling
- Corticosteroid resistant asthma
- Non-adherence and poor inhaler technique
- Concomitant disease and misdiagnosis

Demographic variables
- Age
- Gender
- Race
- Body mass index

Pharmacokinetics of corticosteroids
Table 2 Examples of potential mechanisms of corticosteroids insensitivity in asthma

**Alteration in glucocorticoid receptor (GR) subtypes**

Reduction in ligand-activated GRα subtype numbers or activity

Over expression of the non-functional GRβ subtype

**Defective glucocorticoid receptor binding and nuclear translocation**

Reduction in GR binding affinity and nuclear translocation

**Increase in proinflammatory transcription factors**

Activation of pro-inflammatory transcription factors such as nuclear factor-κB (NF-κB), activator protein-1 (AP-1) and signal transduction-activated transcription factors (STAT)

**Defective histone acetylation**

Reduction in histone deacetylase (HDAC) 2 activity

**Other immune mechanisms**

Suppression of the anti-inflammatory cytokine interleukin (IL)-10 production
Table 3 Current management of corticosteroid insensitivity in asthma

Non-adherence
Employ strategies to improve adherence

Non-pharmacological interventions
Smoking cessation
Weight reduction ‡
Dietary supplementation with vitamin D3 ‡

Current pharmacological treatments

Inhaled corticosteroids
Step-up in dose of inhaled corticosteroid
Extrafine-particle inhaled corticosteroid ‡

Combination of a long acting β₂-agonists (LABA) and an inhaled corticosteroid

Addition of low dose theophylline to an inhaled corticosteroid ††

Leukotriene-receptor antagonist ††

Oral corticosteroid sparing therapies †††
Biological agents e.g. omalizumab, mepolizumab
Immunosuppressive agents e.g. methotrexate

Notes
† Not proven to reverse corticosteroid insensitivity in asthma
†† Possibly in smokers with asthma
††† Unclear whether the reduction in daily oral corticosteroids is due to a reversal of corticosteroid insensitivity or due to the intervention targeting corticosteroid insensitive inflammatory pathways
Table 4 Examples of potential future therapies for corticosteroid insensitivity

**Small molecule drugs**

**Macrolides**

*Protein kinase inhibitors*
- p38 Mitogen-activated protein kinase (MAPK) inhibitors
- Narrow spectrum kinase inhibitors
- Tyrosine kinase inhibitors

*Phosphoinositide 3 (PI3)-kinase inhibitors*
- PI3K-δ inhibitors
- Dual PI3Kδ/γ inhibitors

*Nuclear factor κB (NF-κB) inhibitors*
- IκB kinases (IKKs) inhibitors

*Phosphodiesterase (PDE) inhibitors*
- PDE4 inhibitors
- Dual PDE3 and PDE4 inhibitors

*Peroxisome proliferator-activated receptor-γ (PPARγ) agonists*

**Statins**

*Drugs targeting neutrophilic inflammation*
- C-X-C chemokine receptor (CXCR)2 antagonists
- 5-lipoxygenase-activating protein (FLAP) inhibitors

**Biological agents**

- Interleukin (IL)-17A receptor blockers
- IL-17A blockers
- Tumour necrosis factor (TNF)-α receptor blockers
FIGURE LEGENDS

Figure 1 Variation in therapeutic response to oral corticosteroids in adults with asthma

Distribution of treatment response for change in morning peak expiratory flow (PEF) after a short course of oral corticosteroid in 784 adults with asthma. Reprinted with permission from [5].

Figure 2 Association of serum endotoxin and therapeutic response to oral corticosteroids in adults with asthma

Association of percentage change in FEV$_1$ after oral corticosteroids with sputum endotoxin concentrations in asthmatic patients. The linear regression line (solid line) with the true 95% CI (dashed lines) of nonsmokers (open circles) and smokers (red squares) is shown. In the combined asthma group the percentage change in FEV$_1$ after oral dexamethasone decreased with increasing sputum endotoxin concentration (Spearman rho: $r = -0.329$ [95% CI, 20.570 to 20.036], $p=0.029$). Reprinted with permission from [48].

Figure 3 Addition of low-dose theophylline to inhaled betametasone in smokers with asthma

An exploratory study examined the impact of low dose theophylline added to inhaled corticosteroid (betametasone, BDP) compared to inhaled corticosteroid (BDP) alone in a group of smokers with asthma. Change in peak expiratory flow (PEF) from randomization to 28 days of treatment. Adapted from [109].

Figure 4 Sputum mediator concentrations after atorvastatin and atorvastatin plus inhaled beclometasone or beclometasone alone in smokers with asthma

Sputum mediator concentrations after atorvastatin treatment or placebo and atorvastatin plus inhaled beclometasone or beclometasone alone in smokers with asthma. Examples of
sputum cytokine (IL-1β, G-CSF), chemokine (CCL7) or growth factor (FGF) concentrations after atorvastatin treatment or placebo and atorvastatin plus inhaled beclometasone or beclometasone alone.

Abbreviations: CCL: chemokine (C-C motif) ligand; FGF: Fibroblast growth factor; G-CSF: Granulocyte colony stimulating factor; IL: Interleukin; NS: not significant. Horizontal bar indicates Median (IQR). Adapted from [151].
Figure 1

Change from Run-in After Oral Corticosteroids
Figure 2

A scatter plot showing the relationship between sputum endotoxin log EU/ml and change in FEV1 post corticosteroid %.
**Figure 3**

![Graph showing changes in PEF (l/min) over duration (days) for different treatment groups.]

- **Inhaled BDP**
- Low dose theophylline
- Low dose theophylline + inhaled BDP

*p = 0.008

* p value: BDP + theophylline versus BDP alone
Figure 4

IL-1β

G-CSF

CCL7

FGF2

URL: https://mc.manuscriptcentral.com/errx   Email: Joseph.Walsh@informa.com
RE: Permission to use figure from ERJ

Kay Sharpe [Kay.Sharpe@ersj.org.uk] on behalf of permissions [permissions@ersj.org.uk]

Sent: 27 November 2015 10:48
To: Neil Thomson

Effect of low-dose theophylline plus beclometasone on lung function in smokers with asthma: a pilot study
European Respiratory Journal May 2009, 33 (5) 1010-1017; DOI: 10.1183/09031936.00158208

Material: Figure 2

European Respiratory Society hereby grants you permission to reproduce the material as requested and full acknowledgement must be given.

Acknowledgement Wording: This material has not been reviewed by European Respiratory Society prior to release; therefore the European Respiratory Society may not be responsible for any errors, omissions or inaccuracies, or for any consequences arising there from, in the content. Reproduced with permission of the European Respiratory Society ©: European Respiratory Journal May 2009, 33 (5) 1010-1017; DOI: 10.1183/09031936.00158208

Title: Addressing corticosteroid in adults with asthma
Publication: Expert Review Respiratory Medicine
NLM Title Abbreviation: Expert Rev Respir Med
ISO Abbreviation: Expert Rev Respir Med
Author: Neil C Thomson
Publisher: Taylor & Francis
Publication Format: Print and electronic
1747-6348 (Print)
1747-6356 (Electronic)
1747-6348 (Linking)

Terms & Conditions:

Copyright remains with European Respiratory Society©. These publications are copyrighted material and must not be copied, reproduced, transferred, distributed, leased, licensed, placed in a storage retrieval system or publicly performed or used in any way except as specifically permitted in writing by the publishers (European Respiratory Society), as allowed under the terms and conditions of which it was purchased or as strictly permitted by applicable copyright law.

Reproduction of this is material is confined to the purpose and/or media for which permission is hereby given. Altering/Modifying Material: This is not permitted, however figures and illustrations maybe altered/adapted minimally to serve your work. Please be aware that the permission fee for the requested use of this material is waived in this instance but please be advised that your future requests for materials may attract a fee. This agreement is personal to you and may not be sublicense, assigned or transferred by you to any other person without our written permission.

URL: https://mc.manuscriptcentral.com/errx Email: Joseph.Walsh@informa.com
Any unauthorised distribution or use of this text may be a direct infringement of the publisher’s rights and those responsible may be liable in law accordingly.

This permission is granted for non-exclusive English world rights only.

Regards,

Kay

Kay Sharpe | European Respiratory Society | Publications Office | 442 Glossop Road | Sheffield | S10 2PX | UK
Main Tel: +44 1142672860 | Direct Tel: +44 1142672861 | Fax: +44 1142665064 | E-mail: kay.sharpe@ersj.org.uk