INTRODUCTION.

Approximately 7% of Americans and 2-5% of Russians currently have asthma [1,2,3]. Asthma is characterized by airflow obstruction, bronchial hyperresponsiveness, and inflammation. This inflammation causes recurrent episodes of wheezing, breathlessness, chest tightness, and coughing, which is usually worse at night and in the early morning. There are several inflammatory mediators that have been implicated in the inflammatory process, and management of these mediators can be used to treat the symptoms of asthma. The initiation of asthma therapy is based on the severity of the individual’s asthma [4]. Symptoms are controlled with beta-2-agonists and inhaled corticosteroids in the majority of asthmatics. However many patients with severe asthma and poorly controlled symptoms require additional forms of therapy. This review focuses on current asthma therapy as well as current asthma therapy as well.
emerging novel modalities such as phosphodiesterase inhibitors, Anti-IgE therapy, specific immunotherapy, and bronchothermoplasty.

ICS: Side effects of long term inhaled corticosteroids (ICS)

Asthma is a chronic inflammatory disease with T-helper 2 cell activation. Inhaled corticosteroids are the first-line treatment in adults and children with persistent asthma [5]. There are six different ICS that are available for clinical use: beclomethasone dipropionate, budesonide, flunisolide, fluticasone propionate, mometasone fumarate, and triamcinolone acetate. Beclomethasone disproportionate and Ciclesonide are pro-drugs which are metabolized to active metabolites. Ciclesonide is only converted to its active form in the lower respiratory tract, which helps to reduce local adverse effects such as oral thrush. Low to moderate dosage of ICS is usually adequate for asthma control. However 25—30% of asthma patients with risk of exacerbations need to take high doses to achieve control of their disease [6].

In general, ICS are very well tolerated and have much less concerns of significant side effects compared with systemic corticosteroids. Local adverse effects follow the dual deposition of inhaled corticosteroids on the oropharynx and larynx include oropharyngeal candidiasis, dysphonia, and occasionally coughing from upper airway irritation. The frequency of local adverse effects depends on the dose, frequency and delivery system. For example, patients using MDI may have more complaints of dysphonia than patients using dry powder inhaler. Usually local adverse effects are easy to treat or improve. However, the systemic side effects are of concern when ICS are used for long term, in infants, children and the elderly. It is hard to assess accurately the systemic side effects of ICS because it is usually confounded by disease itself, intermittent use of oral or intravenous corticosteroids, and other co-morbidities. The majority of corticosteroids delivered by MDI are absorbed through gastrointestinal tract and inactivated in liver. About 10—20% of inhaled corticosteroids enter the systemic circulation through the lung parenchyma. With high dose ICS therapy, there are limited data supporting the presence of adrenal suppression, early onset and progression of diabetes [7]. There are no convincing data regarding effects of ICS on bone density, ocular pressure, cataract and respiratory tract infection.

The administration of exogenous corticosteroids results in a negative feedback effect on glucocorticoid receptors in the anterior pituitary gland and hypothalamus, which in turn suppresses levels of corticotropin-releasing hormone and corticotropin, respectively, and a consequent reduction in cortisol secretion from the adrenal cortex. There is no evidence of clinical significant adrenal suppression with long term use of ICS but subclinical adrenal suppression does exist based on bioassays. Measurement of urine cortisol level or early morning plasma cortisol level is used to assess adrenal insufficiency. A meta-analysis of 21 studies of urinary cortisol levels and 13 studies of suppression of 8 AM plasma cortisol levels revealed fluticasone to exhibit significantly steeper dose-related systemic bioactivity than beclomethasone, budesonide, or triamcinolone. These effects were most apparent at doses above 0.8 mg/day [8]. On the other side, treatment with moderate and high doses of ciclesonide does not result in hypothalamus-adrenal axis suppression as compared with placebo [9].

A cohort study by Samy Suissa et al included 388,584 new users of ICS and assessed whether the use and dose of ICS increased the risk of diabetes onset and progression. Use of inhaled corticosteroids was associated with a 34% increase in the rate of diabetes and in the rate of diabetes progression. The risk was greatest with the highest inhaled corticosteroid doses, equivalent to fluticasone 1000 µg per day or more [10].

Physicians prescribing ICS should be aware of the presence of subclinical adrenal suppression and increased rate and progression of diabetes associated with high dose of ICS, especially if the patients are taking a higher than recommended dosage for long term. The physician should evaluate the appropriate dosage of ICS at each visit and attempt to minimize the dosage to obtain asthma control.

LABA: Safety of LABA

There are mainly two forms of LABA available in the market, Formoterol and Salmeterol. Afroformoterol is the R,R-enantiomer of racemic formoterol. Formoterol and salmeterol have similar pharmacological properties: both are highly selective and potent beta₂ agonists. Formoterol has a more rapid onset of action than salmeterol, which makes formoterol suitable for symptom relief as well as symptom prevention in the management of asthma. The duration of action of beta₂ agonists depends on their diffusion microkinetics in the cell membrane lipid bilayer (plasmalemma) and in the aqueous biophase closest to the active site of the beta₂-adrenoceptor. Albuterol is hydrophilic so it does not penetrate the lipid bilayer and it is quickly cleaned by microcirculation. Formoterol and Salmeterol are lipophilic, especial Salmeterol, so they bind to lipid bilayer and have persistent bronchodilator effects. Diffusion theory explains the peak effect observed in Formoterol but not in Salmeterol, and also explains there is a dose dependent response in Formoterol but not in Salmeterol [11].

In general, LABA is recommended when asthma is not controlled with low to moderate dose of ICS. In 2003, the RELIEF trial studied the safety and effectiveness of as needed formoterol compared to salbutamol. Children and adults (n=18,124) were randomized to 6 months as-needed treatment with open-label formoterol 4.5 mg Turbuhaler1 or salbutamol 200 mg pressurized metered dose inhaler or equivalent. This study demonstrates that formoterol as-needed has a similar safety profile to salbutamol, and its use as a reliever therapy is associated with fewer asthma symptoms and exacerbations [12]. However, post-market survey revealed there was possible increased asthma related mortality with regular use of salmeterol. This was further revealed in SMART asthma study. The SMART trial was a multicenter, double-blinded, randomized controlled observation study for 28 weeks. The study was terminated early due to the increased respiratory and asthma related mortality in asthma patient with use of salmeterol. Between 1996 and 2003, 26,355 adult asthma patients were randomized to salmeterol or salbutamol. African Americans represented 18% of the population of patients and the increased respiratory and asthma related mortality was higher in the African American population compared to Caucasians. Whether the greater risk in African Americans reflects genetic predisposition, risk associated with long-acting beta₂ agonist monotherapy, or health maintenance behaviors cannot be determined definitively at this time because this study was not powered or initially designed to study this [13]. Post hoc analysis revealed concurrent use of ICS might have protective effect. Only 38% of African
As recommended by Asthma EPR-3, LABA is added to asthma therapy when asthma is not well controlled with low to moderate dosage of ICS. With the available data, LABA monotherapy for asthma should be rigorously avoided, and combination therapy should be mandatory only when indicated. The mechanism of LABA related severe adverse respiratory events is still unclear. Better understanding of the pharmacokinetics of LABA in asthma will help us identify the appropriate subgroup of patients benefiting from LABA therapy.

Anti-IgE therapy: 10-year clinical experience with Omalizumab

About 5—10% of asthma patient are classified as severe asthma, those who are constantly symptomatic and require frequent bronchodilator therapy with increased rate of asthma exacerbation and asthma-related mortality. More than half of these patients have positive skin test to common allergens. IgE plays a central role in allergic inflammation. IgE binding to the high affinity receptor, FccRI receptors on mast cell and basophils, causes activation of mast cells and basophils, which subsequently release multiple mediators, such as Histamine, LTC4, PGD2 and PAF. In addition, IgE can elevate the FcεRI receptor level on mast cell and basophils and enhance the survival of mast cell. Omalizumab is the only approved anti-IgE therapy for allergic asthma. Omalizumab is a recombinant DNA-derived humanized monoclonal IgG1k antibody binding to Cε3 domain of IgE and forming complexes. This complex prevents IgE from binding to its receptor. Omalizumab was approved for asthma therapy by FDA in June 2003.

Omalizumab can be given subcutaneously or intravenously, but only subcutaneous injection was approved by FDA in U.S.A. The dosage of Omalizumab is calculated based on the weight and pre-treatment IgE level. It can be given every 2 to 4 weeks depending on the dosage. The biowaivability of S.C. Omalizumab is 62%, and the half-life varies from 1 to 4 weeks. It is recommended for patient older than age 12 with moderate to severe persistent asthma, positive skin test or in vitro reactivity to perennial aeroallergens and non-controlled asthma with moderate to high dose ICS. In general, Omalizumab is well tolerated and there was rare anaphylaxis reaction reported (1 to 2 per 1000). There is no adequate safety data for patient younger than 12. IgE less than 30 IU or high than 700 IU, or body weight more than 150 Kg.

Omalizumab reduces circulating levels of IgE in atopic patients to low levels commonly seen in non-atopic individuals, and protects against allergen-induced bronchoconstriction, reduces the need for short acting inhaled β2-agonist and corticosteroids among asthmatic patients. A Cochrane review about the efficacy of omalizumab on ICS usage and asthma exacerbation was released in 2008. In this review, 14 double blinded trials and 3134 moderate to severe allergic asthmatic patients were involved. It confirmed that Omalizumab can reduce the dosage of ICS in moderate and severe asthma, but can only reduce asthma exacerbation in patient with moderate severe asthma. There was no effect on the FEV1 and morning peak flow rate although there was significant effect on quality of life. In further analysis, a significant placebo effect was identified in reducing ICS dosage. In placebo group, 56% had reduction of ICS more than 50% versus 76.7% in Omalizumab therapy group [OR=2.5, 95% CI, (2.02, 3.10)] [18].
Overall, multiple clinical trials proved that Omalizumab can reduce asthma exacerbation in patient with moderate severe asthma, and it is well tolerated. On the other hand, one has to realize the significant placebo effect and high cost associated with Omalizumab therapy ($2706.54/ 4 weeks with maximal dosage of Omalizumab vs. $213.93/ 30 days with Advair 500/50 μg). Patient’s response to anti-IgE therapy is heterogeneous. About one third of the recipients showed substantial improvement, but another third showed no response. With current knowledge, it is difficult to differentiate the responders from nonresponders until after 16-week therapy. Meanwhile, anti-IgE therapy does not completely abrogate high-affinity receptor activation; has a relatively slow onset of efficacy; and, due to dosing limitations, is not approved for patients with very high IgE levels, who might benefit the most from neutralization of serum IgE. Thus, approaches that inhibit high-affinity receptor activation more directly, potently, and quickly than anti-IgE therapy are promising new therapies for the treatment of asthma [19].

PDE4 inhibitor: the indication and role of PDE4 inhibitor in asthma therapy

Cyclic adenosine monophosphate (cAMP) and cyclic guanosine monophosphate (cGMP) are intracellular signalling molecules implicated in the pathophysiology of asthma. They promote smooth muscle relaxation and inhibit inflammation. A novel approach for therapeutic intervention in asthma is through regulation of the phosphodiesterase activity, which is the only cellular pathway currently known for degradation of cAMP and cGMP. Phosphodiesterase type 4 (PDE4) inhibitors increase intracellular concentrations of cAMP. Cyclic AMP acts via protein kinases, which phosphorylates ion channels and signal proteins. This in turn affects epithelial cells, mast cells, macrophages, lymphocytes and neutrophils. Citrolmist and Rolflumist are approved PDE4 inhibitors as an adjuvant therapy for COPD and asthma. And only Rolflumist was approved for asthma therapy [20].

Rolflumist is approved as 500 μg orally once a day for COPD and asthma. It has an 80% oral bioavailability, not affected by food intake, smoking, or diurnal dosing. Rolflumist and its major active metabolite, N-oxide, have T1/2 of 1.5 h and 12 h, respectively and elimination halflives of 10 h and 20 h, respectively. It does not require dose adjustment in patients with severe renal impairment. Also rolflumist and its N-oxide do not interact with inhaled salbutamol or inhaled budesonide. The major side effects are gastrointestinal disturbances, particularly nausea and emesis as well as headache and weight loss [21].

In a double-blind, placebo-controlled, crossover study by Gauvreau et al, 25 subjects with mild allergic asthma were randomized to rolflumist 500 μg or placebo, once daily for two weeks. The primary outcome was the effect of rolflumist on allergen-induced airway eosinophilia. Allergen challenge was performed on Day 14, and FEV1 was measured until 7 hours post challenge. Methacholine challenge was performed on Days 1 (pre-dose), 13 (24 h pre-allergen), and 15 (24 h post-allergen), and spum induction was performed on Days 1, 13, 14 (7 h post-allergen), and 15. Compared to placebo, Rolflumist significantly inhibited the allergen-induced late phase response through inhibiting allergen-induced sputum eosinophil and neutrophil activation [20].

In a review article by Chung, K.F. several trials were analyzed that studied rolflumist and cilomilast. Rolflumist had a small inhibitory effect on the early response to allergen challenge in patients with mild to moderate asthma, and a bigger effect of inhibiting the late response at a dose of 500 μg daily given for 7–10 days prior to allergen. Both rolflumist and cilomilast attenuate exercise-induced bronchoconstriction. In a 3-month study of patients with asthma, there was a dose-dependent improvement in FEV1 and mean morning peak flow rates, a 16% improvement (400 ml) in FEV1 at 500 μg/day dose versus 11% (260 ml) 100 μg/day dose, which was maintained over a 12-month treatment period. In a comparative study of 50 μg/day of rolflumist with inhaled beclomethasone propionate 200 μg twice daily over 12 weeks, equivalent effects were observed in terms of improvement in FEV1 (0.30 L for rolflumist and 0.37 L with beclomethasone), morning peak expiratory flows, reduction in asthma symptoms and reduction in use of rescue medications. The most frequent side-effect recorded in these studies was nausea [22].

The main criticism of all the studies to date with rolflumist in asthma is that they did not include a true randomized placebo group. Based on available data, rolflumist can be used as an adjuvant therapy to inhaled steroids for patient with mild asthma, but there is insufficient data to support the effectiveness of PDE4 inhibitors in severe persistent asthma. And there is no data to compare PDE4 inhibitors with current standard step-up therapy, such as long acting beta₂-agonists, leukotriene inhibitor or theophylline. In addition, the side effects of gastrointestinal disturbances and headache may limit the use of this drug class. However, a number of strategies are currently being pursued in attempts to improve clinical efficacy and reduce side effects of PDE4 inhibitors, including delivery via the inhaled route, development of nonemetic PDE4 inhibitors, mixed PDE inhibitors, and/or antisense biologicals targeted toward PDE4.

Specific Immunotherapy in allergic asthma: the pattern of use and clinical outcome

Specific immunotherapy (SIT) is a method of reducing sensitivity to a given allergen by repeated administration of a dose of that allergen. The primary objectives of allergen-specific immunotherapy are to decrease the symptoms triggered by allergens such as grass, mite, pet dander. This form of therapy involves the subcutaneous administration of gradually increasing quantities of the patient’s relevant allergens until a dose is reached that is effective in inducing immunologic tolerance to the allergens. A Cochrane review of 88 randomized controlled trials examining the use of allergen-specific immunotherapy in asthma management confirmed its efficacy in reducing asthma symptom scores and medication requirements, and improving airway hyperresponsiveness. There was no consistent effect on lung function. Overall, it would have been necessary to treat four patients (95% CI 3 to 6) with immunotherapy to avoid one requiring increased medication. If 16 patients were treated with immunotherapy, one would be expected to develop
a local adverse reaction. If nine patients were treated with immunotherapy, one would be expected to develop a systemic reaction (of any severity) [25]. Similar benefits were suggested with sublingual immunotherapy although the benefit of therapy isn’t large [26]. SIT should be used in addition to bronchodilators and antihistamines for the maximum benefit. Its clinical effects are not as immediate acting and its benefits are often only appreciated in the long term. It has not been shown to be as effective as a single form of therapy.

Currently, specific immunotherapy is the only identified disease-modifying intervention for allergic disease. When used in appropriately-selected patients, allergen-specific immunotherapy is safe. A study on the frequency of systemic adverse reaction of any level of severity associated with SCIT, revealed 82 out of 693 patients (11.3%) developed systemic adverse reaction. Of 82 patients, 69 developed during the build-up phase, and 13 in the maintenance phase. With respect to reaction time, 47 (57%) of the systemic reaction were immediate (within 30 minutes respect to reaction time), and 35 (43%) were delayed [27]. This form of therapy, however, does carry the risk of anaphylactic reactions and, therefore, should only be prescribed by physicians who are adequately trained in the treatment of allergy. Furthermore, immunotherapy should be administered only by physicians who are equipped to manage life-threatening anaphylaxis.

Using recombinant/engineered allergens, possibly modified by site-directed mutagenesis, represents an alternative approach which is directed at maintaining the immunogenicity of a vaccine while reducing the capacity to bind allergen-specific IgE. The results related to their use, hold promise that recombinant allergen–based immunotherapy will improve current immunotherapy practice and may open possibilities for prophylactic vaccination, although no clinical efficacy has been documented yet.

**Bronchithermoplasty: effects on asthma control and clinical evidence**

Many of the symptoms of asthma are thought to be due to smooth muscle contraction, which then leads to bronchoconstriction. Increased airway smooth muscle mass is often found in severe asthma. Bronchial thermoplasty (BT) is performed with Alair Bronchial Thermoplasty System (Asthmatx, Inc; USA), which delivers a controlled amount of thermal energy to the airway through a specific catheter. It decreases airway muscle mass by coagulating bronchial tissue as well as bronchoconstrict response to stimulators, such as methacholine. It may improve symptomatic control and reduce asthma exacerbation [28]. Three controlled clinical trials sponsored by Asthmatrix Inc, were performed to evaluate the efficacy and safety of BT.

The first multicenter prospective randomized control trial, Asthma Intervention Research trial (AIR), studied 112 patients with moderate to severe asthma, measuring a primary outcome of the change in the rate of mild exacerbations between baseline and post treatment. The subjects were between the ages of 18 to 65 and required both inhaled corticosteroids and a long acting beta₂-agonist for control of asthma symptoms. In the intervention group, subjects received three BTs at three week intervals. The clinical outcomes at one year showed BT reduced the frequency of mild exacerbations at a rate equivalent to 10 exacerbations per subject per year and provided 86 additional symptom-free days per subject per year, but it did not decrease the frequency of severe exacerbation. Most frequent adverse effects associated with BT were cough, dyspnea and wheezing, and the majority of the adverse events occurred within 1 day after the procedure and resolved in an average of 7 days after the onset of the event [28].

The second multicenter prospective randomized trial, Research in Severe Asthma (RISA), assessed the safety and efficacy of BT in 32 patients with symptomatic, severe asthma. Similar to the findings in AIR trial, there was a significant increase of short-term asthma related mobility in BT group, including seven hospitalizations [29].

In a multicenter, randomized double blind trial by Castro et al in 2009, also known as the AIR 2 trial, 288 adults were randomized to BT or sham control. In this trial, rather than choosing asthma exacerbation as primary outcome, the authors chose the Asthma Quality of Life Questionnaire (AQLQ). This questionnaire evaluates asthma based on symptoms, limitations of activity, emotional function and environmental factors. Subjects were required at baseline to have an AQLQ score less than 6.25 with higher scores correlating with better quality of life.

In this study, 190 subjects were randomized to the BT group and 98 were randomized to the sham control group. All subjects were scheduled to have three bronchoscopy procedures, 3 weeks apart. BT was administered to the treatment group using the Alair system. In the sham control group the subjects underwent bronchoscopy procedures that were designed to look and sound similar to the initial BT procedure. The subjects were assessed according to the AQLQ along with physical exam, review of symptoms, exacerbations and adverse effects. The subjects were followed up over a 12-month period. A greater improvement in AQLQ was noted in the treatment group comparing to the control group. On the other hand, majority of the patient in the BT group were able to guess correctly that they received the therapy, but not the sham control group. This will influence the AQLQ. There were also fewer exacerbations and ED visits associated with BT [30].

Long term safety of BT was recently reported by Thompson et al in 2011. Patients from the AIR1 trial were followed for adverse events post treatment study. In this study, 45 out of the 52 subjects from the treatment group were studied for 5 years while 24 out of 49 subjects of the control group were studied only for 3 years. It showed that the rate of respiratory adverse events was stable in years 2 through 5 after treatment with BT. The amount of hospitalizations and emergency room visits did not increase. There was no deterioration in FVC and FEV1. The long term benefits from BT were not studied [31].

BT was approved by Food and Drug Administration (FDA) in April 2010. The mechanism of BT has not been fully elucidated, and the long term benefits and adverse effects are still unclear. Even the limited benefit on asthma symptomatic control at one year should be balanced against the short-term adverse effects and cost of procedure. BT should be performed in an experienced center and reserved for patients who demonstrate inability to improve asthma symptoms despite the optimal use of ICS and LABA.

**Conclusion**

Inhaled corticosteroids and long acting beta₂-agonists remain the mainstay of asthma therapy. When asthma symptoms persist despite medication compliance and good inhalational technique, other novel therapies have been shown to be promising. Anti IgE therapy can reduce asthma


