Mechanisms of adverse effects of β-agonists in asthma

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Short-acting β-agonist (SABA) drugs have been mainstays of asthma therapy for many decades and are recommended treatment at all levels of asthma severity, as they provide prompt relief of asthma symptoms through smooth muscle relaxation and, thereby, bronchodilatation. At all levels of asthma severity more severe than mild intermittent, SABAs are recommended to be taken as required for relief of symptoms in conjunction with inhaled corticosteroids (ICSs) taken as regular maintenance treatment. However, in mild asthma SABAs are recommended as monotherapy without concomitant ICS therapy, and in both mild and more severe asthma, greatly increased SABA use at times of asthma exacerbation is almost universal. Here we discuss the safety of inhaled β-agonist monotherapy in asthma and argue against the continued use of β-agonist monotherapy (both short and long acting) in the absence of concomitant ICS therapy in a combination inhaler.

Several epidemiological studies link overuse of SABA therapies at times of asthma exacerbation with increased risk of hospitalisation or mortality. The mechanisms underlying this increased risk have not been clearly determined, but are most likely to involve complex mechanisms including delays in seeking medical care, potential cardiac (tachycardia) and metabolic (hypokalaemia) adverse effects as well as possible effects on underlying asthma severity. Although fenoterol, the SABA linked with the epidemic of asthma mortality in the early 1980s in New Zealand and some other countries, has greater cardiac effects than other SABAs, the reduction in hospitalisations due to asthma exacerbations (along with a reduction in asthma mortality) following the withdrawal of high dose fenoterol in New Zealand in 1990 suggested that the reduction in asthma mortality was not wholly due to reduction in cardiac/metabolic side effects, but probably also due to an effect on disease severity (because if the reduction in mortality were due to reduction in cardiac side effects, the rate of hospitalisations due to asthma exacerbations should have remained unchanged).
The long-acting β-agonists (LABAs) were introduced as asthma treatments in the 1990s, and their use was shown to result in improved lung function and quality of life. Concerns about possible risks associated with LABA treatment were raised very soon after their introduction—these centred around the potential of regular LABAs to reduce bronchodilator sensitivity to β-agonists, to induce tolerance to β-agonists, the risk that the reduction in symptoms achieved with LABAs might reduce awareness of asthma activity and therefore lead to a reduction in usage of ICS therapy or reduced awareness of worsening of asthma control (masking) and finally that if SABAs do have potential to increase disease severity, LABAs might possess this property as well. These concerns were increased by the Salmeterol Multicenter Asthma Research Trial (SMART)—this placebo-controlled study on the safety of the LABA salmeterol in adults with unstable asthma showed a statistically significant, fourfold increase in asthma-related deaths with salmeterol. It was not possible to determine if the risk extended to patients receiving concomitant ICS treatment, nor therefore whether the risk was restricted to the use of salmeterol as monotherapy. In view of these concerns, the Food and Drug Administration (FDA) required black-box warnings on the product labels of both salmeterol and formoterol and recommended that further research was needed into the nature and magnitude of the risk with LABA treatment.

These data led to three important questions relating to β-agonist treatment in asthma. What is the evidence that SABAs or LABAs may increase the underlying severity of asthma, what mechanisms might be involved and what is the evidence that ICSs may protect against an increase in severity consequent upon SABA or LABA treatment?

Regarding the first two questions, regular use of SABA was shown some time ago significantly to increase peak flow variability and bronchial hyper-reactiveness when compared with as-required use during the run-in period of the study, to increase eosinophilic and mast cell responses to allergen challenge, to induce non-specific bronchial hyper-reactivity and to increase sputum eosinophil counts; however, the mechanisms underlying these potential adverse effects are not known.

A report in this issue of Thorax sheds some new light on a potential mechanism that may be involved in adverse effects of β-agonists (see page 763). Lommatzsch et al show that 14 days monotherapy with salmeterol significantly increased brain-derived neurotrophic factor (BDNF) concentrations in serum and platelets of patients with asthma. This increase was abolished by the addition of the ICS fluticasone to the LABA treatment. The induction of BDNF by LABA and protection from this induction by ICSs were confirmed in vitro and, importantly, changes in BDNF concentrations in serum and platelets in the in vivo study correlated with deterioration in airway hyper-responsiveness (AHR) following LABA monotherapy, while, as previously reported many times, when ICSs were added to the LABA treatment there was a significant improvement in AHR. As there is increasing evidence that BDNF may enhance AHR and eosinophilia in allergic airway inflammation, these findings suggest that increased BDNF production may underlie some of the adverse effects of LABA monotherapy. They also suggest that concomitant ICS therapy may exert protective effects through inhibition of this induction.

These data have striking parallels with our own observations with another proinflammatory mediator implicated in asthma pathogenesis in a different experimental setting, and these two reports combined suggest a common molecular mechanism potentially to explain adverse effects of β-agonists in asthma. We studied the effects of β-agonists and ICSs on rhinovirus induction of interleukin-6 (IL-6) as IL-6 is a proinflammatory cytokine and as rhinovirus infections are the major cause of acute exacerbations of asthma. We observed that both salmeterol and the SABA salbutamol increased rhinovirus-induced IL-6 production; the induction of IL-6 by salmeterol could also be induced directly by cAMP, and by other cAMP-elevating agents. Salmeterol increased rhinovirus-induced IL-6 promoter activation via a mechanism dependent upon a cAMP response element (CRE) in the IL-6 promoter. Both SABAs and LABAs are well known to signal their beneficial effects via increasing intracellular levels of cAMP. These data indicate that potential adverse effects (eg, induction of IL-6) are mediated via exactly the same mechanism (induction of cAMP).

Interestingly, BDNF is also regulated via a CRE in its promoter. Although not studied directly by Lommatzsch and colleagues, these facts strongly suggest that the induction of BDNF by salmeterol is likely to have occurred via cAMP-dependent induction of increased BDNF mRNA transcription via the CRE in the BDNF promoter. How many other molecules with potential adverse effects in asthma might also be induced by cAMP? The answer is not known, but a very recent report indicates that Th17 cells and IL-17 production are both directly induced by cAMP. Since Th17 cells and IL-17 have potent proinflammatory properties, it is likely that induction of these responses by cAMP-inducing agents in acute exacerbations of asthma would further increase airway inflammation. A number of other molecules implicated in asthma pathogenesis have also been shown to be regulated via a CRE in their promoter, including cyclo-oxygenase-2 (COX-2), matrix metalloproteinase-2 (MMP-2), and the mucins MUC8 and MUC5AC. These data suggest that β-agonists, while bronchodilating through smooth muscle relaxation, might at the same time have the potential to increase airway obstruction through increasing mucosal inflammation and oedema while simultaneously clogging the airway with secreted mucins.

A further important recent report suggests that the number of other genes that might be similarly regulated via CREs (and therefore potentially be induced by SABA or LABA treatment), might extend into hundreds and possibly thousands of genes as the CREB response element-binding protein (CREB) was found to occupy ~4000 human gene promoter sites in vivo, including large numbers of transcription factors and other genes with potential proinflammatory activities. These studies were not performed in respiratory tissues/cells; however, the above data combined make it clear that SABAs and LABAs have the potential to induce many other genes in asthmatic lung tissue in addition to IL-6 and BDNF.

In our in vitro studies, the ICS fluticasone alone inhibited rhinovirus induction of IL-6, and the addition of fluticasone to salmeterol at equimolar concentrations suppressed the induction of IL-6 observed with salmeterol, indicating that ICSs can counteract the induction of IL-6 observed with LABAs. Similar observations were made by Lommatzsch et al with respect to BDNF, as fluticasone was also shown to reduce BDNF concentrations when used alone in vitro, and, when combined with salmeterol, to block the induction of BDNF both in vitro and in vivo. Interestingly, each of IL-17, MMP-2, MUC8 and MUC5AC, reported to be induced by cAMP or via a CRE in their promoter in the studies cited above, have independently also been shown to be suppressed by steroids. These data suggest that use of β-agonist/ICSs
monotherapy, now with the added evidence of induction of potentially harmful mediators, as well as a molecular mechanism to explain both this induction and a protective effect of ICSs on this adverse effect, we believe that recommendations to combine a β-agonist/ICS in a single inhaler should now be extended to include SABA usage in the treatment of asthma. Combining the ICS with the SABA would ensure that ICS treatment is increased at times of increased disease activity, as well as potentially protecting against adverse effects of SABA overuse at the time of asthma exacerbation.

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