## ASTHMA

# Anticholinergics in the treatment of children and adults with acute asthma: a systematic review with meta-analysis 

G J Rodrigo, J A Castro-Rodriguez

See end of article for
authors' affiliations

Correspondence to: Dr G J Rodrigo, Departamento de Emergencia, Hospita Central de las FF.AA, Av 8 de octubre 3020,
Montevideo 11600,
Uruguay; gurodrig@ adinet.com.uy

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Background: Current guidelines recommend the use of a combination of inhaled $\beta_{2}$ agonists and anticholinergics, particularly for patients with acute severe or life threatening asthma in the emergency setting. However, this statement is based on a relatively small number of randomised controlled trials and related systematic reviews. A review was undertaken to incorporate the more recent evidence available about the effectiveness of treatment with a combination of $\beta_{2}$ agonists and anticholinergics compared with $\beta_{2}$ agonists alone in the treatment of acute asthma.
Methods: A search was conducted of all randomised controlled trials published before April 2005.
Results: Data from 32 randomised controlled trials ( $\mathrm{n}=3611$ subjects) showed significant reductions in hospital admissions in both children ( $R R=0.73 ; 95 \% \mathrm{Cl} 0.63$ to $0.85, \mathrm{p}=0.0001$ ) and adults ( $R R=0.68$; $95 \% \mathrm{Cl} 0.53$ to $0.86, \mathrm{p}=0.002$ ) treated with inhaled anticholinergic agents. Combined treatment also produced a significant increase in spirometric parameters 60-120 minutes after the last treatment in both children (SMD $=-0.54 ; 95 \% \mathrm{Cl}-0.28$ to $-0.81, \mathrm{p}=0.0001$ ) and adults ( $\mathrm{SMD}=-0.36 ; 95 \% \mathrm{Cl}-0.23$ to $-0.49, p=0.00001$ ).
Conclusions: This review strongly suggests that the addition of multiple doses of inhaled ipratropium bromide to $\beta_{2}$ agonists is indicated as the standard treatment in children, adolescents, and adults with moderate to severe exacerbations of asthma in the emergency setting.

Treatment of acute asthma includes inhaled short acting $\beta_{2}$ agonists, systemic corticosteroids (CCS), and supplemental oxygen. ${ }^{12}$ In addition, current guidelines recommend the use of a combination of $\beta_{2}$ agonists and anticholinergics, particularly for patients with acute severe or life threatening asthma. ${ }^{23}$ This statement is based on a relatively small number of randomised controlled trials and related systematic reviews. ${ }^{4-6}$ However, new studies have since been published. ${ }^{7}$ The aim of this systematic review was to update the evidence on the effectiveness of a combination of inhaled anticholinergics and $\beta_{2}$ agonists compared with $\beta_{2}$ agonists alone for the treatment of children, adolescents, and adults with acute asthma in the emergency department (ED).

## METHODS

## Search strategy and selection criteria

The search was conducted using five search strategies to identify potentially relevant trials. (1) MEDLINE (1966-April 2005), EMBASE (1974-April 2005) and CINAHL (1982-April 2005) databases were searched using the following MeSH, full text and keyword terms: emergency OR acute asthma OR status asthmaticus OR severe asthma OR wheeze, AND anticholinergics OR ipratropium OR oxitropium, OR glycopirrolate. (2) An advanced search of the Cochrane Controlled Trials Register (first quarter 2005) was completed using the above search strategy to identify any additional trials. (3) References from included studies, reviews, and texts were searched for citations. (4) Hand searching of the top 20 respiratory journals was completed. (5) We made inquires to Boehringer Ingelheim regarding other published or unpublished trials supported by the company. Trials published solely in abstract form were excluded.

Included studies met the following criteria: (1) Target population: children ( 18 months to 17 years) and adults ( $\geqslant 18$ years) with acute exacerbations of asthma presenting
to an ED or equivalent care setting. (2) Intervention: single or repeated doses of inhaled anticholinergic agents given in combination with inhaled $\beta_{2}$ agonists compared with inhaled $\beta_{2}$ agonists alone. Studies involving the use of atropine were excluded. (3) Design: randomised and placebo controlled trials without language restriction. (4) Primary outcomes: admission to hospital and spirometric testing (final absolute values or change from baseline 60-120 minutes after the last combined anticholinergic and $\beta_{2}$ agonist inhalation). Because the peak bronchodilator effect after the administration of anticholinergics occurs within l-2 hours, it is reasonable to expect significant improvement during this time. Secondary outcome measures were clinical score, duration of treatment in the ED, respiratory rate, oxygen saturation, heart rate, and side effects.

## Data abstraction and validity assessment

Titles, abstracts, and citations were independently reviewed by two reviewers (GJR and JACR) to assess potential relevance for full review. From the full text, both reviewers independently assessed studies for inclusion based on the criteria for population, intervention, study design and outcomes. Data extraction included the following items: (1) Population: age, sex, number of patients studied, patient demographic data, withdrawals. (2) Intervention: agent, dose, route of delivery, and duration of treatment. (3) Control: concurrent treatments. (4) Outcomes. (5) Design: method of randomisation and allocation concealment. Any disagreement over study inclusion was resolved by consensus. The methodological quality of each trial was evaluated

Abbreviations: CCS, corticosteroids; ED, emergency department; FEV ${ }_{1}$, forced expiratory volume in 1 second; NNT, number of patients needed to treat; PEF, peak expiratory flow; RR, relative risk; SMD, standardised mean difference; WMD, weighted mean difference

Table 1 Characteristics of trials in children included in the review

| Study (year) | Design | Language and country | Jadad score | No (and age) of patients | Mean baseline severity | Dose of $\beta$ agonist | Dose of anticholinergic | $\begin{aligned} & \text { CCS } \\ & \text { use } \end{aligned}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Beck et al (1985) ${ }^{12}$ | R, DB | E, Canada | 3 | 25 (6-17 y) | FEV ${ }_{1}<50 \%$ | $\begin{aligned} & \mathrm{S}, 0.05 \mathrm{mg} / \mathrm{kg} \\ & \mathrm{q} 20 \mathrm{~min} \\ & \mathrm{Neb} \times 6 \end{aligned}$ | IB, $0.25 \mathrm{mg} \mathrm{Neb} \times 1$ | No |
| Cook et al (1985) ${ }^{13}$ | R, DB | E, Australia | 4 | $30(18 \mathrm{~m}-12 \mathrm{y})$ | NR | $\begin{aligned} & \mathrm{F}, 0.125-0.5 \mathrm{ml} \\ & \mathrm{Neb} \times 1 \end{aligned}$ | $1 \mathrm{~B}, 1-2 \mathrm{ml} \mathrm{Neb} \times 1$ | No |
| Reisman et al 1988) ${ }^{14}$ | R, DB | E, Canada | 3 | 24 (5-15 y) | $\mathrm{FEV}_{1}<55 \%$ | $\mathrm{S}, 0.05 \mathrm{mg}$ q20 min $\mathrm{Neb} \times 6$ | IB, $0.25 \mathrm{mg} \mathrm{Neb} \times 3$ | No |
| Watson et al (1988) ${ }^{15}$ | R, DB | E, Canada | 3 | $31(6-17 \mathrm{y}$ ) | FEV ${ }_{1}$ 30-70\% | F, 0.62 mg q 60 min $\mathrm{Neb} \times 2$ | IB, 0.25 mg q 60 min Neb $\times 2$ | Yes |
| Phanichyakam et al (1990) ${ }^{1}$ | R, DB | E, Thailand | 1 | 20 (4-15 y) | NR | $\mathrm{T}, 0.5 \mathrm{mg} \mathrm{MDI} \times 1$ | $1 \mathrm{~B}, 0.04 \mathrm{mg} \mathrm{MDI} \times 1$ | No |
| Peterson et al $(1994)^{17}$ | R, DB | E, Canada | 5 | 163 (5-12 y) | FEV ${ }_{1}<70 \%$ | $\mathrm{S}, 3 \mathrm{mg} \mathrm{q} 45 \mathrm{~min}$ Neb $\times 2$ | IB, 0.25 mg q 45 min $\mathrm{Neb} \times 2$ | Yes |
| Schuh et al (1995) ${ }^{18}$ | R, DB | E, Canada | 5 | 80 (5-17 y) | $\mathrm{FEV}_{1}<50 \%$ | $\mathrm{S}, 0.15 \mathrm{mg} / \mathrm{kg} \mathrm{q} 20 \mathrm{~min}$ $\mathrm{Neb} \times 3$ | IB, $0.25 \mathrm{mg} \mathrm{Neb} \times 1$ <br> or IB 0.25 mg Neb $\times 3$ | No |
| Qureshi et al (1997) ${ }^{19}$ | R, DB | E, USA | 5 | 90 (6-18 y) | $\mathrm{FEV}_{1}<50 \%$ | $\mathrm{S}, 0.15 \mathrm{mg} / \mathrm{kg}$ q 30 min $\mathrm{Neb} \times 3$ | IB, $0.5 \mathrm{mg} \mathrm{Neb} \times 2$ | Yes |
| Calvo et al (1998) ${ }^{20}$ | R, DB | Sp, Chile | 3 | 80 (18-55 y) | PEF $<80 \%$ | $\mathrm{S}, 0.2 \mathrm{mg}$ q 15 min MDI $\times 4$ | IB, 0.04 mg ql 5 min MDI $\times 4$ | Yes |
| Ducharme et al (1998) ${ }^{21}$ | R, DB | E, Canada | 5 | 298 (2-18 y) | Mild to moderate | $\mathrm{S}, 0.07 \mathrm{mg} / \mathrm{kg} \mathrm{q} 30 \mathrm{~min}$ Neb | IB, $0.25 \mathrm{mg} \mathrm{Neb} \times 1$ | Yes |
| Qureshi et al (1998) ${ }^{22}$ | R, DB | E, USA | 5 | 434 (2-18 y) | Moderate to severe | $\mathrm{S}, 2.5-5 \mathrm{mg}$ q 20 min $\mathrm{Neb} \times 3$ | $\mathrm{B}, 0.5 \mathrm{mg}$ q 20 min $\mathrm{Neb} \times 2$ | Yes |
| Zorc et al (1999) ${ }^{23}$ | R, DB | E, USA | 5 | 427 (1-17 y) | Moderate to severe | $\begin{aligned} & \mathrm{S}, 2.5 \mathrm{mg} \mathrm{q} 20 \\ & \mathrm{Neb} \times 3 \end{aligned}$ | $\mathrm{BB}, 0.5 \mathrm{mg}$ q20 min $\mathrm{Neb} \times 3$ | Yes |
| Benito Fernandez et al (2000) ${ }^{24}$ | R, SB | Sp, Spain | 5 | $102(5 \mathrm{~m}-16 \mathrm{y}$ ) | Severe | $\mathrm{S}, 0.2 \mathrm{mg} / \mathrm{kg} 930 \mathrm{~min}$ $\mathrm{Neb} \times 2$ | IB, 0.25 mg q30 min $\mathrm{Neb} \times 2$ | Yes |
| SienraMonge <br> et al $(2000)^{25}$ | R, DB | Sp, Mexico | 2 | 30 (8-15 y) | Moderate to severe | $\mathrm{S}, 0.2 \mathrm{mg} \mathrm{q10} \mathrm{~min}$ MDI $\times 3$ | $\mathrm{IB}, 0.02 \mathrm{mg}$ q10 min MDI $\times 3$ | No |
| Timsit et al (2002) ${ }^{26}$ | R | F, France | 3 | 114 (2-15 y) | Moderate | $\mathrm{S}, 0.15 \mathrm{mg} / \mathrm{kg} \mathrm{q} 20 \mathrm{~min}$ $\mathrm{Neb} \times 6$ | $\mathrm{IB}, 0.25 \mathrm{mg} \mathrm{q} 20 \mathrm{~min}$ $\mathrm{Neb} \times 3$ | Yes |
| Sharma et al (2004) ${ }^{27}$ | R | E, India | 2 | 50 (6-14 y) | Moderate to severe | $\mathrm{S}, 0.15 \mathrm{mg} / \mathrm{kg} \mathrm{q} 20 \mathrm{~min}$ $\mathrm{Neb} \times 3$ | IB, 0.25 mg q 20 min Neb $\times 3$ | No |

R, randomised; SB, single blind; DB, double blind; E, English; Sp, Spanish; F, French; FEV ${ }_{1}$, forced expiratory volume in 1 second; PEF, peak expiratory flow; NR, not reported;
S, salbutamol; F, fenoterol; $T$, terbutaline; IB, ipratropium bromide; CCS, systemic corticosteroids.
using the 5-point scale ( $0=$ worst and $5=$ best) described by Jadad et al. ${ }^{8}$ This instrument assesses the adequacy of randomisation, blinding, and the handling of withdrawals and drop outs.

## Data analysis

The data were combined in the meta-analysis by means of random effects models. ${ }^{9}$ Binary outcomes were pooled using common relative risk ( RR ) and $95 \%$ confidence intervals (CI). The number of patients needed to treat (NNT) to prevent the adverse outcome of interest was calculated. For continuous outcomes the weighted mean difference (WMD) (for variables using the same unit of measure) or the standardised mean differences (SMD) (reported in SD units where different units were used) and $95 \%$ CI were calculated. We tested for heterogeneity using the DerSimonian and Laird Q statistic and also measured heterogeneity with the $\mathrm{I}^{2}$ test. ${ }^{10}$ Values of $25 \%, 50 \%$, and $75 \%$ represent low, moderate, and high heterogeneity, respectively. Publication bias was evaluated by means of formal statistical analysis. ${ }^{11}$ Otherwise, a p value of $<0.05$ using a two tailed test was considered significant. When heterogeneity was found, subgroup analyses were carried out in an attempt to explain the findings. Sensitivity analysis was performed to identify sources of heterogeneity. These subgroups included: intensity of anticholinergic protocol, baseline severity, co-therapies, and methodological quality of the studies. The meta-analysis was performed using Review Manager 4.2.7 software (Cochrane Review Manager, Cochrane Collaboration, Oxford, UK, 2004).

## RESULTS

A total of 88 studies were examined in full text for possible inclusion, 56 of which were excluded for the following reasons: non-randomised trials $(\mathrm{n}=4)$, non-acute asthma ( $\mathrm{n}=14$ ), anticholinergics alone were studied $(n=6)$, hospitalised
patients $(\mathrm{n}=8)$, use of atropine $(\mathrm{n}=5)$, chronic asthma ( $\mathrm{n}=18$ ), and use of intravenous route $(\mathrm{n}=1)$. A total of 32 randomised controlled trials ( 16 including children and adolescents ${ }^{12-27}$ and 16 including adults ${ }^{28-43}$ ) were therefore selected for further analysis (tables 1 and 2). Five studies were supported by Boehringer Ingelheim. ${ }^{17}{ }^{18}{ }^{35-37}$ Data for 3611 subjects ( 1564 children and adolescents, 2047 adults) were available for meta-analysis. There was a total agreement between the two independent reviewers on inclusion of studies and Jadad study quality grading. The anticholinergic agent used was ipratropium bromide in 29 studies, ${ }^{12-32} 34-38404243$ oxitropium bromide in two studies, ${ }^{39}{ }^{41}$ and glycopyrrolate in one study. ${ }^{33}$ Trials were grouped according to the intensity of the anticholinergic treatment: those testing the addition of a single dose of an anticholinergic agent to $\beta_{2}$ agonist inhalations were named single dose protocols, and those testing more than one dose were grouped as multiple dose protocols. Thirteen studies (five in children ${ }^{1213161821}$ and eight in adults ${ }^{28} 29{ }^{31-33}$ 363843 ) tested a single dose protocol and the remaining 19 trials used more than one dose of anticholinergic. Of these, 18 studies tested multiple doses in a predetermined fixed regimen (multiple dose fixed protocol) and one study tested the addition of anticholinergics to every $\beta_{2}$ agonist inhalation, leaving the number of inhalations determined by the patient's needs (multiple dose flexible protocol). ${ }^{20}$ One trial tested the first two protocols. ${ }^{18}$ Asthma severity was defined at baseline by spirometric testing (forced expiratory volume in 1 second $\left(\mathrm{FEV}_{1}\right)$ or peak expiratory flow (PEF) $70-50 \%$ of predicted $=$ moderate exacerbation, and $\mathrm{FEV}_{1}$ or $\mathrm{PEF}<50 \%$ of predicted $=$ severe exacerbation) or different clinical scores. Most enrolled patients with acute asthma had moderate to severe exacerbations, but several studies reported data stratified on asthma severity. 2 2-24 374142 The most frequently reported outcomes were hospital admission (20 studies) and spirometry (26 studies); respiratory resistance measured by forced oscillation was used

Table 2 Characteristics of trials in adults included in the review

| Study (year) | Design | Language and country | Jadad score | No (and age) of patients | Mean baseline severity | Dose of $\beta$ agonist | Dose of anticholinergic | $\begin{aligned} & \text { CCS } \\ & \text { use } \end{aligned}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Bryant (1985) ${ }^{28}$ | R, DB | E, Australia | 2 | 28 ( $\geqslant 18 \mathrm{y}$ ) | FEV ${ }_{1}<75 \%$ | F, $1 \mathrm{mg} \mathrm{Neb} \times 1$ | IB, $0.5 \mathrm{mg} \mathrm{Neb} \times 1$ | No |
| Rebuck et al (1987) ${ }^{29}$ | MC, R, DB | E, Canada | 4 | $148(\geqslant 18 y)$ | $\mathrm{FEV}_{1}<70 \%$ | F, $1.25 \mathrm{mg} \mathrm{Neb} \times 1$ | IB, $0.5 \mathrm{mg} \mathrm{Neb} \times 1$ | Yes |
| Higgins et al (1988) ${ }^{30}$ | R, DB | E, England | 2 | $40(\geqslant 18 y)$ | PEF $<30 \%$ | $\mathrm{S}, 5 \mathrm{mg} \mathrm{q} 120 \mathrm{~min}$ $\mathrm{Neb} \times 2$ | $\mathrm{IB}, 0.5 \mathrm{mg}$ q 120 min Neb $\times 2$ | Yes |
| O'Driscoll et al (1989) ${ }^{31}$ | R, DB | E, England | 2 | $56(\geqslant 18 y)$ | PEF $<35 \%$ | S, $10 \mathrm{mg} \mathrm{Neb} \times 1$ | IB, $0.5 \mathrm{mg} \mathrm{Neb} \times 1$ | Yes |
| Summers and Tarala (1990) ${ }^{32}$ | $R, ~ D B$ | E, Australia | 3 | 76 (16-70 y) | PEF $<60 \%$ | S, $5 \mathrm{mg} \mathrm{Neb} \times 1$ | IB, $0.5 \mathrm{mg} \mathrm{Neb} \times 1$ | Yes |
| Cydulka and Emerman (1994) ${ }^{33}$ | R, DB | E, USA | 3 | 125 ( $\geqslant 18 y$ ) | $\mathrm{FEV}_{1}<75 \%$ | $\mathrm{S}, 2.5 \mathrm{mg} \mathrm{q} 60 \mathrm{~min}$ $\mathrm{Neb} \times 3$ | Gly, $2 \mathrm{mg} \mathrm{Neb} \times 1$ | Yes |
| Rodrigo and Rodrigo $(1995)^{34}$ | R, DB | Sp, Uruguay | 3 | 22 (18-50 y) | $\mathrm{FEV}_{1}<50 \%$ | $\mathrm{S}, 0.4 \mathrm{mg} \mathrm{q} 10 \mathrm{~min}$ MDI $\times 3 \mathrm{~h}$ | $\mathrm{IB}, 0.08 \mathrm{mg} \mathrm{q} 10 \mathrm{~min}$ MDI $\times 3 \mathrm{~h}$ | No |
| Karpel et al (1996) ${ }^{35}$ | MC, R, DB | E, USA | 5 | 384 (18-55 y) | $\mathrm{FEV}_{1}<60 \%$ | $\mathrm{S}, 2.5 \mathrm{mg} \mathrm{q} 45 \mathrm{~min}$ $\mathrm{Neb} \times 2$ | IB, 0.5 mg q 45 min Neb $\times 2$ | No |
| FitzGerald et al $(1997)^{36}$ | MC, R, DB | E, Canada | 3 | 342 (18-50 y) | $\mathrm{FEV}_{1}<70 \%$ | $\mathrm{S}, 3 \mathrm{mg} \mathrm{Neb} \times 1$ | IB, $0.5 \mathrm{mg} \mathrm{Neb} \times 1$ | Yes |
| Garret et al (1997) ${ }^{37}$ | TC, R, DB | E, New Zealand | 4 | 338 (18-55 y) | $\mathrm{FEV}_{1}<70 \%$ | $\mathrm{S}, 2.5 \mathrm{mg} \mathrm{q} 45 \mathrm{~min}$ Neb $\times 2$ | $\mathrm{IB}, 0.5 \mathrm{mg} \mathrm{q} 45 \mathrm{~min}$ Neb $\times 2$ | Yes |
| Lin et al (1998) ${ }^{38}$ | R, DB | E, USA | 4 | 55 ( $\geqslant 18 \mathrm{y}$ ) | PEF $<200 \mathrm{l} / \mathrm{min}$ | $\mathrm{S}, 2.5 \mathrm{mg}$ q 20 min $\mathrm{Neb} \times 3$ | IB, $0.5 \mathrm{mg} \mathrm{Neb} \times 1$ | No |
| Kamei et al (1999) ${ }^{39}$ | MC, R | E, Japan | 3 | 64 ( $\geqslant 18 \mathrm{y}$ ) | $\mathrm{FEV}_{1}<70 \%$ | F, 0.2 mg q 1 min MDI $\times 5$ | $O B, 0.1 \mathrm{mg} \mathrm{ql} \mathrm{min}$ MDI $\times 5$ | Yes |
| Weber et al (1999) ${ }^{40}$ | $R, ~ D B$ | E, USA | 5 | 67 ( $\geqslant 18 \mathrm{y}$ ) | PEF $<70 \%$ | $\begin{aligned} & \mathrm{S}, 10 \mathrm{mg} \text { ql } \mathrm{h} \\ & \mathrm{Neb} \times 3 \mathrm{~h} \end{aligned}$ | IB, 1 mg ql h Neb $\times 3 \mathrm{~h}$ | No |
| Nakano et al (2000) ${ }^{41}$ | R, SB | E, Japan | 4 | 74 ( $\geqslant 18 \mathrm{y}$ ) | PEF $<50 \%$ | $\mathrm{S}, 0.4 \mathrm{mg} \mathrm{q} 20 \mathrm{~min}$ MDI $\times 3$ | $\mathrm{OB}, 0.4 \mathrm{mg}$ q 20 min MDI $\times 3$ | Yes |
| Rodrigo and Rodrigo (2000) ${ }^{42}$ | R, DB | E, Uruguay | 5 | 180 (18-50 y) | $\mathrm{FEV}_{1}<50 \%$ | $\mathrm{S}, 0.4 \mathrm{mg} \mathrm{q} 10 \mathrm{~min}$ MDI $\times 3 \mathrm{~h}$ | IB, 0.08 mg q 10 min MDI $\times 3 \mathrm{~h}$ | No |
| Aggarwal et al $(2002)^{43}$ | R | E, India | 2 | 48 (13-50 y) | PEF < $50 \%$ | $\mathrm{S}, 5 \mathrm{mg} \mathrm{q} 60 \mathrm{~min}$ Neb $\times 2$ | IB, $0.5 \mathrm{mg} \mathrm{Neb} \times 1$ | No |

in one trial. ${ }^{21}$ One study did not provide spirometric data or admission rates. ${ }^{13}$ Clinical scores were used in only a few studies and the reporting of adverse effects was variable.

## Hospital admissions

Ten studies accumulating 1786 children and adolescents reported hospital admissions. ${ }^{14} 17-19{ }^{21-24} 2627$ One study tested two protocols (single and multiple fixed dose) ${ }^{18}$ and three trials reported data stratified by asthma severity (moderate and severe patients). ${ }^{22-24}$ At the end of treatment patients
who received inhaled $\beta_{2}$ agonists and anticholinergics had a significantly lower admission rate (fig l). The NNT was 13 ( $95 \%$ CI 9 to 28 ), indicating that 13 children needed to be treated with $\beta_{2}$ agonists and anticholinergics to prevent one admission. There was no evidence of systematic bias identified by the measure of funnel plot asymmetry. Also, no significant heterogeneity was demonstrated, which accepts the null hypothesis of similar treatment effects. Stratification on the basis of baseline severity (moderate $v$ severe) and the intensity of the anticholinergic protocol


Figure 1 Pooled relative risk for hospital admission (with $95 \%$ confidence interval) of eligible studies in children comparing the addition of anticholinergic agents to $\beta_{2}$ agonists (treatment) with $\beta_{2}$ agonists alone (control). Trials stratified according to intensity of anticholinergic treatment (single or multiple fixed dose protocols) and asthma severity (moderate or severe patients).
(single $v$ multiple fixed dose protocol) suggested a trend towards a reduced risk of admission in children with the most severe asthma attack and treated with multiple doses of anticholinergics. The NNT to prevent one admission among severe patients was 7 ( $95 \%$ CI 4 to 16). The hospital admission rate did not change when we excluded studies without explicit admission criteria ( $\mathrm{RR}=0.73 ; 95 \%$ CI 0.62 to $\left.0.85, \mathrm{I}^{2}=0 \%\right)$. The use of systemic CCS did not modify this outcome ( $\mathrm{RR}=0.69 ; 95 \%$ CI 0.58 to 0.81 ).

Nine trials totalling 1556 adults with acute asthma reported hospital admissions. ${ }^{33-38} 40-42$ One trial reported data stratified on asthma severity (moderate and severe patients). ${ }^{42}$ There was a significant reduction in the hospital admission rate favouring anticholinergic use (fig 2). The NNT was 14 ( $95 \%$ CI 9 to 30 ). There was no evidence of systematic bias identified by the measure of funnel plot asymmetry. Again, no significant heterogeneity was demonstrated. Stratification on the basis of baseline severity (moderate $v$ severe) and the intensity of the anticholinergic protocol (single $v$ multiple fixed dose) suggested a trend towards a reduced risk of admission in adults with the most severe asthma attack and treated with multiple doses of anticholinergics (fig 2). Intensity of anticholinergic treatment greatly influenced the reduction in hospital admission; a greater reduction was seen in trials using three or more doses of anticholinergic agents $(\mathrm{RR}=0.53 ; 95 \%$ CI 0.36 to 0.76 , $\mathrm{p}=0.0006$; NNT $=6$; $95 \%$ CI 4 to 13 ). These results did not change when only studies with explicit admission criteria were pooled ( $\mathrm{RR}=0.58$; $95 \%$ CI 0.38 to $0.87, \mathrm{I}^{2}=28 \%$ ) or when systemic CCS were used ( $\mathrm{RR}=0.74 ; 95 \%$ CI 0.48 to 1.14 ).

## Spirometric testing

Nine studies examined the response to treatment in children and adolescents with acute asthma using spirometry. ${ }^{12} 14-161819222527$ Five trials reported the percentage change in $\mathrm{FEV}_{1},{ }^{12}{ }^{14-1625}$ three reported the percentage change in PEFR, ${ }^{1922} 27$ one reported the change in percentage predicted $\mathrm{FEV}_{1},{ }^{18}$ and one study reported the percentage change in respiratory resistance. ${ }^{21}$ One trial tested two protocols ${ }^{18}$ (single and multiple fixed dose) and one study presented data
stratified by severity of obstruction (moderate and severe). ${ }^{22}$ Data were recorded 60-120 minutes after the last combined treatment. When all the studies were pooled a significant improvement in spirometric parameters favoured the combination treatment $(\mathrm{SMD}=-0.54 ; 95 \% \mathrm{CI}-0.28$ to -0.81 , $\mathrm{p}=0.0001)$. However, there was significant heterogeneity $\left(\chi^{2}=23.41, \mathrm{df}=10, \mathrm{I}^{2}=57.3 \%, \mathrm{p}=0.009\right)$. When we pooled the seven studies that reported $\mathrm{FEV}_{1}$ data (change in percentage predict or percentage change) ${ }^{12} 14-16182527$ stratified by the intensity of anticholinergic treatment (one or two doses $v$ more than two doses), homogeneity was achieved (fig 3). The use of more than two doses of anticholinergics showed more benefit than lower doses. There was no evidence of systematic bias. Patients treated with one or two doses of anticholinergic agents had a mean difference of change in $\mathrm{FEV}_{1}$ of $12.4 \%$ ( $95 \%$ CI 5.4 to 19.4) compared with those who did not receive anticholinergics, while those who received more than two doses had a mean difference of $16.3 \%$ ( $95 \%$ CI 8.2 to 24.5).

Spirometric data were reported by 16 studies in adult subjects. ${ }^{28-43}$ Two trials showed data stratified by severity of obstruction (moderate and severity). ${ }^{41}{ }^{42}$ Eight trials reported $\mathrm{FEV}_{1}(\mathrm{l}),^{28} 29{ }^{34-37} 39{ }^{42} 11$ reported PEFR ( $\mathrm{l} / \mathrm{min}$ ), , ${ }^{29-322^{34} 38-43}$ and one reported $\mathrm{FEV}_{1}$ (\% predicted). ${ }^{33}$ Combined treatment produced a significantly greater increase in spirometric parameters than $\beta_{2}$ agonists alone $(\mathrm{SMD}=-0.36 ; 95 \% \mathrm{CI}-0.23$ to -0.49 , $\mathrm{p}=0.00001$ ). There was a significant heterogeneity between trials $\left(\chi^{2}=25.5, \mathrm{df}=15, \mathrm{I}^{2}=41.3 \%, \mathrm{p}=0.04\right)$. Homogeneity was achieved when studies that reported PEFR (l/min) were stratified by intensity of anticholinergic treatment (fig 4). ${ }^{29-32} 3438-43$ Again, the use of more than two doses of anticholinergics produced a greater benefit than one or two doses and there was no evidence of systematic bias. As previously observed for PEFR, patients treated with more than two doses of anticholinergics had a significant difference in $\mathrm{FEV}_{1}$ of 0.44 l ( $95 \%$ CI 0.25 to 0.63 ) while those treated with one or two doses had a difference of only 0.15 l ( $95 \%$ CI 0.05 to 0.24 ).

## Other outcomes

Three paediatric studies ${ }^{21} 22{ }^{24}$ reported a significant reduction in the clinical score after combined treatment ( $\mathrm{SMD}=-0.29$;


Figure 2 Pooled relative risk for hospital admission (with $95 \%$ confidence interval) of eligible studies in adults comparing the addition of anticholinergic agents to $\beta_{2}$ agonists (treatment) with $\beta_{2}$ agonists alone (control). Trials stratified according to intensity of anticholinergic treatment (single or multiple fixed dose protocols) and asthma severity (moderate or severe patients).


Figure 3 Pooled standardised mean difference (with $95 \%$ confidence interval) in forced expiratory volume in the first second (change in percentage predicted or percentage change) of children studies comparing the addition of anticholinergic agents to $\beta_{2}$ agonists (treatment) with $\beta_{2}$ agonists alone (control). Trials stratified according to the intensity of anticholinergic treatment (one or two doses $v$ more than two doses).
$95 \%$ CI -0.51 to $-0.07, \mathrm{p}=0.01)$. No significant heterogeneity was demonstrated $\left(\chi^{2}=1.33, \mathrm{df}=3, \mathrm{p}=0.72\right.$, $\mathrm{I}^{2}=0 \%$ ). There was no apparent increase in the occurrence of side effects among subjects treated with either single or multiple dose protocols. Thus, there was no significant difference between groups in the five studies in children that reported the presence of tremor $(\mathrm{RR}=1.15 ; 95 \%$ CI 0.79 to $1.69, \mathrm{p}=0.46$ ). ${ }^{1314171827}$ An identical pattern was seen in three adult studies that reported the same variable $(R R=1.28 ; 95 \%$ CI $0.92-1.78, p=0.14) .{ }^{31} 3542$ Six adult trials that evaluated the effect of treatment on heart rate did not find a difference between groups ( $\mathrm{WMD}=-2.07$; 95\% CI -4.35 to $0.21, \mathrm{p}=0.07) .{ }^{28} 2931343542$ There was insufficient information to pool outcomes such as oxygen saturation due to the insufficient number of trials reporting this outcome. Analysis of the only trial which tested the administration of multiple inhalations of combined treatment until a satisfactory clinical response was achieved (multiple dose flexible protocol) showed a significant decrease in the clinical score at 30-45 minutes between patients treated with salbutamol and ipratropium and those treated with salbutamol alone. ${ }^{20}$

## DISCUSSION

This systematic review constitutes an effort to incorporate the best evidence available up to April 2005 on the role of inhaled anticholinergic agents added to $\beta_{2}$ agonists in children,
adolescents, and adults with acute asthma in the ED setting New data were found which we added to previous review. ${ }^{35}$ Thus, 10 new randomised trials (four in children ${ }^{24-27}$ and six in adults ${ }^{33} 34^{39}{ }^{41-43}$ ) with a total of 809 patients have been added, representing an increase of $22 \%$ on the previous sample. Unlike the previous reviews, this study has enabled analysis of the effect of cumulative doses, particularly in adult studies. Several important conclusions can be made. Overall, our analysis confirmed that early administration of inhaled anticholinergic agents with $\beta_{2}$ agonists lead to a reduction in admission rates of both children and adults of $30 \%$. Baseline severity and the intensity of the anticholinergic protocol clearly influenced the magnitude of the benefit. Thus, anticholinergic agents are particularly beneficial in patients with moderate to severe obstruction (FEV ${ }_{1}<70 \%$ of predicted) treated with multiple dose fixed protocols consisting of three or more doses of an anticholinergic. These patients had a reduction in the hospital admission rate of 30$45 \%$ and only $6-14$ subjects need to be treated to prevent one hospital admission. This is a very relevant finding since hospital admissions count for the largest part of direct health costs for asthma in most countries, and children or adults with more severe asthma attacks are more prone to be admitted to hospital. However, this review did not identify any beneficial effects of anticholinergic agents in patients with mild acute asthma. The fact that the use of systemic CCS


Figure 4 Pooled weighted mean difference (with $95 \%$ confidence interval) in peak expiratory flow (l/min) of studies in adults comparing the addition of anticholinergics to $\beta_{2}$ agonists (treatment) with $\beta_{2}$ agonists alone (control). Trials were stratified by intensity of anticholinergic treatment (one or two doses $v$ more than two doses).
has not shown a significant effect is in agreement with the evidence that they require 6-12 hours to modify outcomes such as hospital admission or spirometric parameters. ${ }^{44}{ }^{45}$ The short duration of the study period in all trials made it unlikely that these drugs would have a significant contribution.

Significant differences favouring the combination treatment were observed on spirometric data in both children and adults. Again, there was a dose-response relationship, with a greater benefit being achieved in patients treated with more than two doses of anticholinergic agents in combination with a $\beta_{2}$ agonist. In adults, treatment with more than two doses produced clinically significant improvements in both $\mathrm{FEV}_{1}$ ( 0.44 l ) and PEFR ( $50.5 \mathrm{l} / \mathrm{min}$ ). ${ }^{46}$

In our meta-analysis we also looked at secondary outcomes and side effects but these were difficult to analyse because there was insufficient information to be pooled. A few of the studies in children reported a significant reduction in different clinical scores after combined treatment. There was no apparent increase in the occurrence of side effects such as tremor or heart rate among subjects treated with single or multiple dose protocols.

## Strengths and limitations of the study

This study met most of the methodological criteria suggested for scientific reviews. ${ }^{47}$ Similar to all systematic reviews, this meta-analysis is limited by the quality and quantity of existing research and how data are reported. A comprehensive search of the published literature for potentially relevant studies was conducted using a systematic strategy to avoid bias. All of the 32 trials were randomised, and 26 were double blind. Exclusion of trials with lower methodological quality did not affect the conclusions. Assessment of the consistency of effects across studies is an essential part of the review to determine the generalisability of the findings; low values of heterogeneity ( $<15 \%$ ) were obtained in all group and subgroup comparisons. The generalisability of study results to different countries should also be considered, particularly with regard to the hospital admission criteria. The decision to admit patients is based on many factors including past asthma and current exacerbation histories and spirometric test results, as well as clinical factors. Important variations in admission criteria could therefore influence the results. However, the results did not change when we analysed only studies with explicit criteria for admission to hospital.

## Authors' affiliations

G J Rodrigo, Departamento de Emergencia, Hospital Central de las FF.AA, Av 8 de Octubre 3050, Montevideo 11600 and Clinica Respirar, Benito Nardone 2310, Montevideo 11300, Uruguay J A Castro-Rodriguez, Pediatric Pulmonary Section, Department of Pediatrics, School of Medicine, University of Chile, Santiago, Chile
GJR has received fees for speaking from Boehringer Ingelheim.

## REFERENCES

1 Rodrigo GJ, Rodrigo C, Hall JB. Acute asthma in adults. A review. Chest 2004;125:1091-102.
2 National Institutes of Health. Global strategy for asthma management and prevention, NIH Publication 02-3659, 2004.
3 British Thoracic Society/Scottish Intercollegiate Guidelines Network. British guideline on the management of asthma: management of acute asthma. guideline on the management of
Thorax 2003;58(Suppl I):i32-50.
4 Rodrigo G, Rodrigo C, Burschtin O. A meta-analysis of the effects of ipratropium bromide in adults with acute asthma. Am J Med 1999;107:363-70.
5 Stoodley RG, Aaron SD, Dales RE. The role of ipratropium bromide in the emergency management of acute asthma exacerbation: a meta-analysis of randomized clinical trials. Ann Emerg Med 1999;34:8-18.
6 Plotnick LH, Ducharme FM. Combined inhaled anticholinergics and beta2agonists for initial treatment of acute asthma in children. Cochrane Database of Systematic Reviews, 2000, Issue 3, CD000060.
7 Rodrigo GJ, Rodrigo C. The role of anticholinergics in acute asthma treatment. An evidence-based evaluation. Chest 2002;121:1977-87.

8 Jadad AR, Moore RA, Carroll D, et al. Assessing the quality of reports of randomized clinical trials: is blinding necessary? Control Clin Trials 1996;17:1-12.
9 DerSomonian R, Laird N. Meta-analysis in clinical trials. Control Clin Trials 1986;7:177-88.
10 Higgins JPT, Thompson SG, Deecks JJ, et al. Measuring inconsistency in metaanalyses. BMJ 2003;327:557-60.
11 Egger M, Davey Smith G, Schneider M, et al. Bias in meta-analysis detected by a simple, graphical test. BMJ 1997;315:629-34.
12 Beck R, Robertson C, Galdes-Sebaldt $M$, et al. Combined salbutamol and ipratropium bromide by inhalation in the treatment of severe acute asthma. J Pediatr 1985;107:605-68.
13 Cook JJ, Fergusson DM, Dawson KP. Ipratropium and fenoterol in the treatment of acute asthma. Pharmatherapeutica 1985;4:383-6.
14 Reisman J, Galdes-Sebaldt M, Kazim F, et al. Frequent administration of inhalation of salbutamol and ipratropium bromide in the initial management of severe acute asthma children. J Allergy Clin Immunol 1988;81:16-20.
15 Watson WTA, Becker AB, Simmons FER. Comparison of ipratropium solution, fenoterol solution and their combination administered by nebulizer and face mask to children with acute asthma. J Allergy Clin Immunol 1988;82:1012-8.
16 Phanichyakam P, Kraisarin C, Sasisakulporn C. Comparison of inhaled terbutaline and inhaled terbutaline plus ipratropium bromide in acute asthmatic children. Asian Pacific J Allergy Immunol 1990;8:45-8.
17 Peterson R, Wensley D, Mitchell I, et al. Boehringer Ingelheim Trial No. 2442430, 3. 1994.
18 Schuh S, Johnson DW, Callahan S, et al. Efficacy of frequent nebulized ipratropium added to frequent high-dose albuterol therapy in severe childhood asthma. J Pediatr 1995;126:639-45.
19 Qureshi FA, Zaritsky A, Lakkis H. Efficacy of nebulized ipratropium in severe asthmatic children. Ann Emerg Med 1997;29:205-11.
20 Calvo GM, Calvo AM, Marin HF, et al. Is it useful to add an anticholinergic treatment to beta2-adrenergic medication in acute asthma attack? J Invest Allergy Clin Immunol 1998;8:30-4.
21 Ducharme FM, Davis GM. Randomized controlled trial of ipratropium bromide and frequent low doses of salbutamol in the treatment of mild and moderate acute asthma. J Pediatr 1998;133:479-85.
22 Qureshi F, Pestian J, Davis P, et al. Effect of nebulized ipratropium on the hospitalization rates of children with asthma. N Engl J Med hospitalization rates
23 Zorc JJ, Pusic MV, Ogborn CJ, et al. Ipratropium bromide added to asthma treatment in the pediatric emergency department. Pediatrics 1999;103:748-52.
24 Benito Fernández J, Maintegui Raso S, Sánchez Echanitz J, et al. Eficacia de la administración precoz de bromuro de ipratropio nebulizado en niños con crisis asmática. An Esp Pediatr 2000;53:217-22.
25 Sienra Monje JJL, Bermejo Guevara MA, del Rio Navarro BE, et al. Grado y duración de la broncodilatación mediante la administración de un agonista $\beta 2$ solo vs un agonista $\beta 2$ mas bromuro de ipratropio en niños con asma aguda. Rev Alergia México 2000;XLVII:26-9.
26 Timsit S, Sannier N, Bocquet N, et al. Apport du bromure d'ipratropium dans la prise en charge des crises d'asthme aux urgencies. Arch Pédiatr 2002;9:117-24.
27 Sharma A, Madaan A. Nebulized salbutamol vs salbutamol and ipratropium combination in asthma. Indian J Pediatr 2004;71:121-4.
28 Bryant DH. Nebulized ipratropium bromide in the treatment of acute asthma. Chest 1985;88:24-9.
29 Rebuck AS, Chapman KR, Abboud R, et al. Nebulized anticholinergic and sympathomimetic treatment of asthma and chronic obstructive airways disease in the emergency room. Am J Med 1987;82:59-64.
30 Higgins RM, Stradling JR, Lane DJ. Should ipratropium be added to betaagonists in treatment of acute severe asthma? Chest 1988;94:718-22.
31 O'Driscoll BR, Taylor RJ, Horsley MG, et al. Nebulised salbutamol with and without ipratropium bromide in acute airflow obstruction. Lancet 1989;i:1418-20.
32 Summers QA, Tarala RA. Nebulized ipratropium in the treatment of acute asthma. Chest 1990;97:430-4.
33 Cydulka RK, Emerman CL. Effects of combined treatment with glycopyrrolate and albuterol in acute exacerbation of asthma. Ann Emerg Med 1994;23:270-4.
34 Rodrigo G, Rodrigo C. Tratamiento de la crisis asmática con altas dosis de salbutamol y bromuro de ipratropio administrados mediante inhalador de dosis medida e inhalocámara. Pac Critico 1995;8:175-84.
35 Karpel JP, Schacter EN, Fanta Ch, et al. A comparison of ipratropium and albuterol vs. albuterol alone for the treatment of acute asthma. Chest 1996;110:611-6.
36 FitzGerald JM, Grunfeld A, Pare PD, et al. The clinical efficacy of combination nebulized anticholinergic and adrenergic bronchodilators vs. nebulized adrenergic bronchodilator alone in acute asthma. Chest 1997;111:311-5.
37 Garret JE, Town GI, Rodwell P, et al. Nebulized salbutamol with and without ipratropium bromide in the treatment of acute asthma. J Allergy Clin Immunol 1997;100:165-70.
38 Lin RY, Pesola GR, Bakalchuk L, et al. Superiority of ipratropium plus albuterol over albuterol alone in the emergency department management of adult asthma: a randomized clinical trial. Ann Emerg Med 1998;31:208-13.
39 Kamei T, Fujita J, Okada H, et al. Comparison between fenoterol and fenoterol plus oxytropium bromide delivered by metered-dose inhaler with inspirease to relief acute asthma attack. J Asthma 1999;36:67-75.
40 Weber EJ, Levitt A, Covington JK, et al. E. Effect of continuously nebulized ipratropium bromide plus albuterol on emergency department length of stay
and hospital admission rates in patients with acute bronchospasm. A randomized, controlled trial. Chest 1999;115:937-44.
41 Nakano Y, Enomoto N, Kawamoto A, et al. Efficacy of adding multiple doses of oxitropium bromide to salbutamol delivered by means of a metered-dose inhaler with a spacer device in adults with acute severe asthma. J Allergy Clin Immunol 2000;106:472-8.
42 Rodrigo GJ, Rodrigo C. First-line therapy for adult patients with acute asthma receiving a multiple-dose protocol of ipratropium bromide plus albuterol in the emergency department. Am J Respir Crit Care Med 2000;161:1862-8.
43 Aggarwal P, Singh O, Wali JP, et al. Efficacy of nebulised ipratropium in acute bronchial asthma. J Indian Acad Clin Med 2002;3:353-9.

44 Rodrigo G, Rodrigo C. Corticosteroids in the emergency department therapy of acute adult asthma. An evidence-based evaluation. Chest 1999;116:285-95.
45 Rowe BH, Spooner C, Ducharme FM, et al. Early emergency department treatment of acute asthma with systemic corticosteroids. Cochrane Database of Systematic Reviews, 2001, Issue 1, CD002178.
46 Santanello NC, Zhang J, Seidenberg B, et al. What are minimal important changes for asthma measures in clinical trial? Eur Respir J 1999;14:23-7.
47 Moher D, Cook DJ, Eastwood S, et al. Improving the quality of reports of meta-analyses of randomized controlled trials: the QUOROM statement. Lancet 1999;354:1896-900.

## LUNG ALERT

## Nitric oxide protects against airway hyperresponsiveness

A Que LG, Liu L, Yan Y, et al. Protection from experimental asthma by an endogenous bronchodilator. Science 2005;308:1618-21

N
itric oxide (NO) is a highly active endogenous bronchodilator and, although increased levels are found in asthmatic lungs, the link between NO and asthma has remained elusive. NO is short lived in vivo but it reacts with cysteine sulphurs (thiols) in proteins to form more stable $S$-nitrosothiols (SNOs) which act as a source of bioactive NO. $S$-nitrosoglutathione (GSNO) is the most abundant SNO found in the airways where its levels are governed by the enzyme GSNO reductase (GSNOR). However, GSNO is depleted in asthmatic airways, suggesting a protective role.
In this study the authors showed that GSNOR levels were raised in the lungs of mice exposed to the allergen ovalbumin (OVA), probably due to lysis of airway epithelial cells and leucocytes. SNO levels were depleted. GSNOR gene knockout mice exposed to OVA had raised levels of SNOs in the airway, reduced basal airway tone, and no response to methacholine. Levels of type II inducible NO synthase were similar to wild type mice, as was the inflammatory response measured by bronchoalveolar fluid cell counts and IL-13, serum total IgE, and mucus metaplasia. Tracheal rings from wild type mice became desensitised to repeated $\beta$ adrenergic stimulation, whereas GSNOR knockout mice did not and so retained the capacity to relax.
This is the first study to show a definitive link between NO and airway hyperresponsiveness (AHR). NO, when present as SNOs, protects against AHR through modulation of $\beta$ adrenoreceptor function. SNO levels are regulated by GSNOR which is raised in asthmatic airways, and the resulting lack of SNOs promotes AHR.

P Kewin
Wellcome Clinical Research Fellow, Department of Respiratory Medicine, Gartnavel General Hospital Glasgow, UK; pk49y@clinmed.gla.ac.uk

