ocs

Evidence Table 11. Pharmacologic Therapy: Inhaled Corticosteroids—Combination Therapy

Abbreviations used in table:

AE	adverse event	OR	odds ratio
AM	morning	PEF	peak expiratory flow
ECP	eosinophil cationic protein	RR	relative risk
FEV ₁	forced expiratory volume in 1 sec	SABA	short-acting beta2-agonist
ICS	inhaled corticosteroid	SAE	serious adverse event
ITT	intent-to-treat	SMD	standardized mean difference
LABA	long-acting beta ₂ -agonist	95% CI	95 percent confidence interval
LTRA	leukotriene receptor antagonist	WMD	weighted mean difference

oral corticosteroids

^{*} indicates primary outcome

Evidence Table 11. Pharmacologic Therapy: Inhaled Corticosteroids—Combination Therapy

		Study Population			
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)	
ICS + LABA vs. ICS					
Lemanske et al. Inhaled corticosteroid reduction and elimination in patients with persistent asthma receiving salmeterol. JAMA 2001;285(20):2594–2603. (National Heart, Lung, and Blood Institute)	Multicenter, randomized, controlled, triple-blinded, double-dummy, parallel group (6 National Institutes of Health-sponsored, university-based ambulatory care centers)	175 (167)	(at end of salmeterol introduction phase) Age 12–65 yr, mean = 35 yr Gender 49% male, 51% female Ethnicity 64% White, 22% Black, 8% Hispanic, 6% other	(at end of salmeterol introduction phase) Persistent asthma FEV ₁ mean = 2.5 L before salmeterol FEV ₁ % pred. mean = 73.6 before salmeterol Morning PEF mean = 431.2 L/min Evening PEF mean = 437.5 L/min PEF variability mean = 0.12	
Ni et al. Addition of inhaled long-acting beta ₂ -agonists to inhaled steroids as first line therapy for persistent asthma in steroid-naive adults. Cochrane Database Syst Rev 2005;(2): CD005307.	Meta-analysis of randomized controlled trials; 4 rated as high quality, 3 as moderate quality, and 1 as poor quality	8 trials with 1,061 subjects (1 trial had 2 control and 2 intervention groups and was counted as 2 trials, resulting in 9 trials for the meta-analysis).	Age Mean ranged from 12–77 yr. Note: None of the 5 trials indicated the proportions who were 12–18 years of age. Gender Males ranged from 25% to 61%.	Mild (5 trials) to moderate (4 trials) airway obstruction Mean FEV ₁ % pred. ranged from 66% to 96%. Naïve to both LABA and ICS	
Ni et al. Long-acting beta ₂ -agonists versus placebo in addition to inhaled corticosteroids in children and adults with chronic asthma. Cochrane Database Syst Rev 2005;(4):CD0055355. (Canadian Cochrane Network, McGill University, Canada)	Meta-analysis of randomized controlled trials; 24 were parallel group designs and 2 were crossover studies. (23 of the 26 trials rated of high quality using Jadad scale.)	26 trials with 31 comparisons; sample sizes ranged from 16 to 663, with 8,147 total participants.	Age In 18 adult trials, mean ages ranged from 35 to 48 yr; in 8 pediatric studies, mean age ranged from 8.5 to 14 yr. Gender Males ranged from 30% to 71%.	Moderate (17 trials) or mild (8 trials); unreported in 1 trial In 23 trials, participants had inadequate control; in 3 trials, participants were asymptomatic and well-controlled. FEV₁ % pred. mean ranged from 51% to 79% in 17 trials; mean was ≥80% in 8 trials and unreported in 1 trial.	

		Study Population			
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)	
Weiler et al. Effect of fluticasone/salmeterol administered via a single device on exercise-induced bronchospasm in patients with persistent asthma. Ann Allergy Asthma Immunol 2005;94(1):65–72. (GlaxoSmithKline)	Multicenter, randomized, double-blind, parallel-group design (40 sites in US)	192 (ITT analysis)	Age 12–50 yr, mean = 29 yr Gender 39% male, 61% female Ethnicity 72% White, 20% Black, 5% Hispanic, 3% other Smoking 85% had never used tobacco. None had smoked in past year. None had more than 10 pack-years of smoking.	Persistent asthma History of asthma ≥15 yr, 59% FEV₁ mean = 2.72L FEV₁ % pred. mean = 78 Maximal postexercise decline in FEV₁, mean = 31.6 Asthma-related emergency care in previous year, 15% Asthma-related hospitalization in previous year, 5% 82% were using fluticasone before enrollment.	
ICS + LTRA vs. ICS					
Robinson et al. Addition of Randomized, double-blind, 10		100 (72)	Age 22–79 yr, mean = 52.3 yr Gender 38% male, 62% female Ethnicity Not reported Smoking 3% current smokers, 20% exsmokers, 67% never smokers	Moderate or severe asthma $ FEV_1 \ \% \ pred. \ median = 59.7 $ $ FVC \ \% \ pred., \ median = 89.9 $ $ PEF \ \% \ pred., \ 63.6 $ $ 23\% \ had \ some \ irreversible \ airflow \ obstruction. $ $ All \ were \ taking \ ICS: \ 12\% \ beclomethasone \ propionate, \ 23\% \ budesonide, \ 54\% \ fluticasone \ propionate. $	

		Study Population			
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)	
Simons et al. Montelukast added to budesonide in children with persistent asthma: a randomized, double-blind, crossover study. J Pediatr 2001; 128(5):694–698. (Merck & Co., Inc.)	Multicenter, randomized, double-blind, crossover design	279 (251)	Age 5-15 yr, mean = 10.4 yr Gender 67% male, 33% female Ethnicity 83% White, 10% Asian, 6% Hispanic, 1% other Height 109-182 cm, mean = 144 cm	Persistent asthma FEV₁ = 1.8 L FEV₁ % pred. mean = 77.7 FEV₁ reversibility mean = 18.1% FVC % pred. median = 89.9 Morning PEF mean = 315 L/min Mean beta₂-agonist use = 2.9 puffs/day Treated with inhaled glucocorticoid for ≥6 weeks	
Vaquerizo et al. Effect of montelukast added to inhaled budesonide on control of mild to moderate asthma. Thorax 2003; 58(3):204–210. (Merck Sharp & Dohme, Spain)	Multicenter, randomized, double-blind, parallel group design (80 hospital centers in Spain)	639 (573 completed; ITT analysis based on 625) Age 18–79 yr, mean = 43 yr Gender 62% male, 38% female Ethnicity Not reported Smoking All nonsmokers, 33% exsmokers		Mild-to-moderate asthma Duration mean = 13.8 yr FEV ₁ mean = 2.5L FEV ₁ % pred. mean = 81 PEF mean = 369 L/min Daily beta-agonist use, 3.2 puffs/day Budesonide, mcg/day: 400–800, 67% 801–1,200, 5% 1,201–1,600, 27%	
ICS + LABA vs. ICS + LTRA					
Nelson et al. Fluticasone propionate/salmeterol combination provides more effective asthma control than low-dose inhaled corticosteroid plus montelukast. J Allergy Clin Immunol 2000; 106(6):1088–1095. (GlaxoWellcome Inc.)	Multicenter, randomized, double-blind, double-dummy, parallel group (39 sites in United States)	487 (ITT analysis)	Age 15–83 yr, mean = 41.6 yr Gender 39% male, 61% female Ethnicity 87% White, 13% other	Asthma for ≥ 6 months Taking low-to-moderate doses of ICS for ≥ 30 days before screening FEV ₁ , mean = 2.39 L FEV ₁ % pred., mean = 70.4 Morning PEF, mean = 395 L/min Evening PEF, mean = 415 L/min Average use of ≥ 4 puffs/day of albuterol	

		Study Population				
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)		
Fish et al. Salmeterol powder provides significantly better benefit than montelukast in asthmatic patients receiving concomitant inhaled corticosteroid therapy. Chest 2001;120(2): 423–430. (GlaxoWellcome Inc.)	Multicenter, randomized, double-blind, double-dummy, parallel group (2 trials at 71 clinical centers in United States and Puerto Rico)	948 (948)	Age 15–83 yr, mean = 39.7 yr Gender 38.8% male, 61.2% female Ethnicity 85.0% White, 7.5% Black, 5.6% American Hispanic, 1.9% other	Mild-to-moderate persistent asthma for ≥ 6 months Duration, $24.7\% < 10$ yr, $75.3\% \geq 10$ yr FEV ₁ , mean = 2.3 L FEV ₁ % pred., mean = 68.3 Symptomatic >6 weeks prior to screening Constant dosage of ICS for 30 days prior to screening Mean (range) ICS use: 482 mcg (44–1,760) fluticasone, 552 mcg (100–1,600) triamcinolone, 265 mcg (84–672) beclomethasone, 651 mcg (84–1,200) budesonide, 1,077 mcg (250–2,000) flunisolide		
Nelson et al. Comparison of inhaled salmeterol and oral zafirlukast in asthmatic patients using concomitant inhaled corticosteroids. MedGenMed 2001;3(4):3.	Multicenter, randomized, double-blind, double-dummy, parallel-group, clinical trial (data from 2 trials) (54 clinical centers)	429 (429)	Age 12–89 yr, mean = 40 yr Gender 44% male, 56% female Ethnicity 87.9% Caucasian, 6.3% Black, 1.4% Asian, 4.4% American Hispanic	Mild to moderate persistent asthma Diagnosis for ≥6 months Duration of asthma: 24% <10 yr, 76% ≥10 yr FEV₁ before bronchodilation, mean = 2.27 L FEV₁ % pred., 66.2 Stable dose of ICS taken for at least 30 days; dosages were not recorded.		
Bjermer et al. Montelukast and fluticasone compared with salmeterol and fluticasone in protecting against asthma exacerbation in adults: one year, double blind, randomised, comparative trial. BMJ 2003; 327(7420):891. (Merck & Co.)	Multicenter, randomized, double-dummy, double-blind, parallel group (148 sites in 37 countries)	1490 (ITT analysis)	Age 15–72 yr; mean = 41.1 yr Gender 45% male, 55% female Ethnicity 77.6% White, 0.7% Black, 7.1% Asian, 14.6% other	Chronic asthma for ≥ 1 yr $FEV_1 \text{ mean} = 2.4 \text{ L}$ $FEV_1 \% \text{ pred., mean} = 72.0$ $FEV_1 \% \text{ reversibility, } 18.6$ $PEF, \text{ mean} = 386 \text{ L/min}$ $Mean \text{ use of beta}_2\text{-agonist} = 3.3 \text{ puffs/day}$ $Mean \text{ number of nocturnal awakenings} = 2.6 \text{ days/week}$		

		Study Population				
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)		
Ringdal et al. The salmeterol/fluticasone combination is more effective than fluticasone plus oral montelukast in asthma. Respir Med 2003;97(3):234–241. (GlaxoSmithKline)	Multicenter, randomized, double-blind, double-dummy, parallel-group study (114 centers in 19 countries)	806 (805 for safety population; 725 for ITT analysis)	Age 14–79 yr, mean = 43 yr Gender 45% male, 55% female Smoking 6.2% current smoker, 22.6% exsmoker, 71.2% nonsmoker	$FEV_1, mean = 2.60 \ L$ $FEV_1 \% \ pred., mean = 75$ $Reversibility \ mean = 27.2\%$ $Morning \ PEF \ mean = 369 \ L/min$ $All \ received \ ICS \ (400-1,000 \ mcg/day \ of \ beclomethasone \ dipropionate, \ budesonide, \ or \ flunisolide; \ or \ 200-500 \ mcg/day \ of \ fluticasone \ propionate) \ for \ at \ least \ 4 \ weeks \ before \ the \ study.$		
Ceylan et al. Addition of formoterol or montelukast to low-dose budesonide: an efficacy comparison in short- and long-term asthma control. Respiration 2004;71(6):594–601.	Randomized comparison trial			Moderately persistent asthma Persistent symptoms for ≥1 yr Duration of asthma, mean = 8.6 yr History of allergic rhinitis, 65% Use of ICS for ≥6 months Morning PEF, mean = 264.6 FEV ₁ , mean = 2.4 L FEV ₁ % pred., mean = 70.5 Beta ₂ -agonist use, mean = 2.4 puffs/day		
Ilowite et al. Addition of montelukast or salmeterol to fluticasone for protection against asthma attacks: a randomized, double-blind, multicenter study. Ann Allergy Asthma Immunol 2004;92(6):641–648.	Multicenter, randomized, double-dummy, double-blind, parallel group (132 centers in United States)	1,473 (1,452; modified ITT analysis)	Age 14–73 yr, mean = 38.6 yr Gender 39.4% male, 60.6% female Ethnicity 84.1% White, 8.9% Black, 5.2% Hispanic, 1.8% other	Chronic asthma for ≥1 yr FEV₁ % pred., mean = 74.3 FEV₁ % reversibility, 18.6 Mean use of beta₂-agonist = 3.5 puffs/day Mean number of nocturnal awakenings = 1.88 days/week Used beta₂-agonist, on average, once/day during last 14 days of run-in period Used ICSs daily for at least 8 weeks before first study visit		

		Study Population			
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)	
Ram et al. Long-acting beta ₂ -agonists versus anti-leukotrienes as add-on therapy to inhaled corticosteroids for chronic asthma. Cochrane Database of Syst Rev 2005;(1):CD003137. (NHS Research and Development UK) NOTE: Meta-analysis includes studies abstracted separately above: Bjermer et al. 2003; Fish et al. 2001; llowite et al. 2004; Nelson et al. 2000; and Nelson et al.	Meta-analysis of randomized controlled trials, all rated of high quality	8 trials with 5,895 patients	Age All adults; mean ages ranged from 35 to 43	Recurrent or persistent asthma Mean duration of asthma ranged from 10 to 26 years. Moderate airway obstruction ranged from 66% to 76% FEV ₁ predicted. Subjects were symptomatic at enrollment. ICS doses at enrollment were <400–560 mcg/day of beclomethasone or equivalent.	
ICS + LTRA vs. increasing I	CS				
Price et al. A randomized controlled trial of montelukast plus inhaled budesonide versus double dose inhaled budesonide in adult patients with asthma. Thorax 2003;58(3): 211–216. (Merck & Co.)	Multicenter, randomized, double-blind, placebo- controlled, parallel-group noninferiority study	889 (843; ITT analysis)	Age 15–75 yr, mean = 43 yr Gender 40% male, 60% female Ethnicity 76.9% White, 0.7% Black, 4.9% Asian, 17.4% other Smoking All nonsmokers or exsmokers	Duration >6 months, mean = 17 yr FEV_1 mean = 2.3 L FEV_1 % pred., mean = 68.7 PEF , mean = 384 L/min Daily beta ₂ -agonist use, 2.7 puffs/day Nocturnal awakenings, median 13.3% of days Days missed from work/school due to asthma in previous year, mean = 21.5 days 41.4% used OCS in previous year.	

		Study Population			
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)	
ICS + LABA vs. increasing	ics				
Jenkins et al. Salmeterol/fluticasone propionate combination therapy 50/250 microg twice daily is more effective than budesonide 800 microg twice daily in treating moderate to severe asthma. Respir Med 2000;94(7):715–723. (GlaxoWellcome)	Multisided, randomized, double-blind, double-dummy, parallel group (multinational study in 44 centers)	353 (ITT analysis)	Age 14–80 yr, mean = 46 yr Gender 50% male, 50% female Ethnicity Not reported	Moderate-to-severe asthma Duration: 6%. 0 to <1 yr; 18%, 1 to <5 yr; 17%, 5 to <10 yr; 60%, ≥10 yr FEV₁ % pred., 33–109, mean = 70 Corticosteroid therapy: 24% using fluticasone propionate (median 500 mcg/day), 48% using budesonide (median 800 mcg/day), 29% using beclomethasone dipropionate (median 1,000 mcg/day)	
Matz et al. Addition of salmeterol to low-dose fluticasone versus higher-dose fluticasone: an analysis of asthma exacerbations. J Allergy Clin Immunol 2001; 107(5):783–789. (GlaxoWellcome Inc.)	Retrospective analysis of 2 identical, multicenter, randomized, double-dummy, double-blind, parallel-group studies (71 research centers)	925 (ITT analysis)	Age ≥12 yr Gender Not reported Ethnicity Not reported	Persistent asthma	

		Study Population				
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)		
O'Byrne et al. Low dose inhaled budesonide and formoterol in mild persistent asthma: the OPTIMA randomized trial. Am J Respir Crit Care Med 2001;164(8 Pt 1): 1392–1397.	Multicenter, randomized, double-blind, parallel group (198 centers in 17 countries)	Group A: 900 Group B: 1,625	Group A Age ≥12 yr, mean = 30.8 yr Gender 40% male, 60% female Ethnicity Not reported Group B Age ≥12 yr, mean = 37.2 yr Gender 43% male, 57% female Ethnicity Not reported	Group A Mild asthma No ICS for ≥3 mo FEV₁ % pred., mean = 89.7 Morning PEF, mean = 418 L/min Days with symptoms, mean = 39.8% Nights with awakenings, mean = 11.0% Group B Mild asthma ≤400 mcg/day budesonide or its equivalent for ≥3 mo FEV₁ % pred., mean = 86.5 Morning PEF, mean = 4,189 L/min Days with symptoms, mean = 37.8% Nights with awakenings, mean = 6.7%		
Bateman et al. Combination therapy with single inhaler budesonide/formoterol compared with high dose of fluticasone propionate alone in patients with moderate persistent asthma. Am J Respir Med 2003;2(3):275–281. (AstraZeneca)	Multicenter, randomized, double-blind, double-dummy, parallel group (37 centers in 6 countries)	344 (344)	Age 17–75 yr, mean = 42 yr Gender 43% male, 57% female Ethnicity Not reported Smoking 6% smokers, 24% exsmokers, 70% never smoked	Moderate persistent asthma Duration of asthma, ≥6 months; mean = 16.3 yr FEV ₁ , geometric mean = 2.4 L FEV ₁ % pred., mean = 78 Morning PEF, mean = 359 Prestudy ICS dose, mean = 594 mcg/day		

		Study Population				
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)		
Ind et al. Addition of salmeterol to fluticasone propionate treatment in moderate-to-severe asthma. Respir Med 2003;97(5):555–562. (GlaxoWellcome Research & Development)	Multicenter, randomized, double-dummy, double-blind, parallel group (100 hospitals and primary care centers in 6 countries)	502 (496 ITT analysis)	Age ≥16 yr, mean = 44.8 yr Gender 46% male, 54% female Ethnicity Not reported Smoking 17.5% smokers, 33.7% exsmokers, 48.8% never smoked	Moderate-to-severe asthma Duration: 0.2–68 yr FEV ₁ , mean = 2.3 L FEV ₁ % pred., mean = 74.5 PEF % pred., mean = 74.5 Median ICS daily dose 1,000 mcg budesonide/beclomethasone dipropionate (BDP) In the past year, 20.8% required hospitalization, 67% required OCS, and 86.5% required other therapy changes.		
Lalloo et al. Budesonide and formoterol in a single inhaler improves asthma control compared with increasing the dose of corticosteroid in adults with mild-to-moderate asthma. Chest 2003;123(5): 1480–1487. (AstraZeneca, Lund, Sweden)	Multicenter, randomized, double-blind, parallel group (51 centers in 7 countries)	467 (430; ITT analysis)	Age 18–78 yr, mean = 41 yr Gender 43% male, 57% female Ethnicity Not reported	Mild-to-moderate asthma Duration: 6 months–53 yr; mean = 11.5 yr FEV ₁ % pred., 38–157; mean = 81 FEV ₁ % reversibility, 11–98; mean = 22 Dose of inhaled steroid = 200–500 mcg/day; mean = 387 mcg/day		
Bergmann et al. Salmeterol/fluticasone propionate (50/250 microg) combination is superior to double dose fluticasone (500 microg) for the treatment of symptomatic moderate asthma. Swiss Med Wkly 2004; 134(3-4):50–58. (GlaxoWellcome)	Multicenter, randomized, double-blind study (76 private practices or outpatient clinics at hospitals)	365 (347 ITT analysis)	Age ≥18 yr, mean = 49.3 yr Gender 46.4% male, 53.6% female Ethnicity Not reported Smoking Nonsmokers or exsmokers	Moderate asthma Duration of asthma: 33.5%, 1–5 yr; 66.5%. 5–19 yr Start of ICS: 21.6%, 3–5 yr; 78.4%, >5 yr FVC % pred., mean = 87.7 FEV ₁ % pred., mean = 75.1 PEF % pred., 73.3 Morning PEF, mean = 317 Percentage symptom-free days, mean = 16.5 Rescue medications, mean = 2.6 puffs/day		

		Study Population				
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)		
Jonsson et al. An economic evaluation of combination treatment with budesonide and formoterol in patients with mild-to-moderate persistent asthma. Respir Med 2004;98(11): 1146–1154.	Multicenter, randomized, prospective, double-blind, parallel-group trial (17 countries)	1,272 for clinical outcomes; 1,233 for economic analysis	No characteristics provided. See O'Byrne et al. for description of sample.	Mild-to-moderate persistent asthma. See O'Bryne et al. for description.		
Masoli et al. Moderate dose inhaled corticosteroids plus salmeterol versus higher doses of inhaled corticosteroids in symptomatic asthma. Thorax 2005;60(9): 730–734.	Meta-analysis of double-blind, randomized trials	12 studies with 4,576 subjects	Age ≥12 yr Gender Not reported Ethnicity Not reported	Moderate to severe Symptomatic on moderate doses of ICS (200 mcg/day fluticasone or equivalent) FEV ₁ % pred., mean = 64%; range = 40%–85% across studies		
LABA A vs. LABA B in addi	tion to ICS					
Palmqvist et al. Onset of bronchodilation of budesonide/formoterol vs. salmeterol/fluticasone in single inhalers. Pulm Pharmacol Ther 2001;14(1):29–34.	Randomized, double-blind, placebo-controlled crossover study	30 (30)	Age 28–73 yr, mean = 49 yr Gender 50% male, 50% female	FEV ₁ , range 1.48–4.28 L. mean = 2.54 L FEV ₁ % pred., range 60.6–98.6. mean = 78.2 Reversibility after 0.1 mg salbutamol, range 5–22%, mean = 12% Reversibility after 0.5 mg salbutamol, range 12–31%, mean = 19% All used inhaled SABAs as needed; 50% used LABAs as needed.		
Everden et al. Eformoterol Turbohaler compared with salmeterol by dry powder inhaler in asthmatic children not controlled on inhaled corticosteroids. Pediatr Allergy Immunol 2004;15(1):40–47.	Multicenter, randomized, open, parallel-group comparative study (56 general practice centers in the United Kingdom and 2 in the Republic of Ireland)	156 (155)	Age 6-17 yr, mean = 11.7 yr 52% 6-11 yr, 48% 12-17 yr Gender 57% male, 43% female Height Mean = 148 cm Weight Mean = 46 kg	Moderate persistent asthma Constant dose of ICS ≥4 weeks prior to enrollment; range, 100–1,600 mcg/day, mean = 362 mcg/day Run-in use of SABA: mean = 0.84 inhalations/night; mean = 1.16 inhalations/day at school; mean = 3.95 inhalations/24 hours PEF mean = 314.5 L/min		

	Stu	Study Characteristics				Findings			
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety		
ICS + LABA vs. ICS									
Lemanske et al. Inhaled corticosteroid reduction and	Purpose/Objective: To determine vin patients with persistent asthma after								
elimination in patients with persistent asthma receiving salmeterol. JAMA 2001;285(20): 2594–2603. (National Heart, Lung, and Blood Institute)	Arm 1 Triamcinolone + salmeterol xinafoate (n=154; 148 completed and randomized to Arms 1a and 1b) Arm 1a Triamcinolone + salmeterol (S+) (n=74); 71 completed and continued with triamcinolone + salmeterol (69 completed) Arm 1b Triamcinolone + salmeterol (S-) (n=74); 71 completed and assigned to placebo triamcinolone + salmeterol) (n=66 completed)	400/42 mcg twice daily 400/42 mcg twice daily 200/42 mcg twice daily	2-week salmeterol introduction phase, 8-week triamcinolone reduction phase, and 8-week triamcinolone elimination phase after a 6-week run-in period Albuterol used for rescue therapy as needed. Randomization at Phase II was by ethnic group, sex, and age.			8.3% (95% CI 2% to 15%) in the S– group. At end of elimination phase, treatment failure occurred in 46.3% (95% CI 34% to 59%) of S– group and 13.7% (95% CI 5% to 22%) of S+ group. RR of treatment failure during reduction phase for S– vs. S+ was 2.2 (95% CI 0.5 to 9.2, p = 0.27; Cox regression model), and during the elimination phase, RR was 4.3 (95% CI 2.0 to 9.2, p <0.001).			
	Arm 2 Triamcinolone + placebo salmeterol (n=21); 19 completed and assigned to triamcinolone + placebo salmeterol (P-); 18 completed and assigned to placebo triamcinolone + placebo salmeterol	42 mcg twice daily salmeterol 400 mcg twice daily triamcinolone placebo							

	Stu	dy Characteristics		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
Ni et al. Addition of inhaled long-acting beta ₂ -agonists to inhaled steroids as first line therapy for persistent asthma in steroid-naive adults. Cochrane Database of Syst Rev 2005;(2):CD005307.	ICS		Duration was 4–8 weeks (4 trials), 12 weeks (2 trials), 24 weeks (1 trial), and 52 weeks (2 trials).	Difference favored LABA for improvement from baseline in FEV ₁ in standard deviation units (SMD 0.29, 95% CI 0.17 to 0.42; 0.42, p <0.00001; 6 trials), in mL (WMD 210 mL, 95% CI 120 to 300; 5 trials), and in morning PEF (WMD 21.4 L/min, 95% CI 15.36 to 27.45, p <0.0001; 5 trials). No difference in change in PEF variability occurred (SMD -0.04, 95% CI -0.50 to 0.41; 4 trials).		*No difference occurred in risk of exacerbation requiring systemic corticosteroids (RR 1.19, 95% CI 0.75 to 1.88; 3 trials). Reduction in symptom score occurred for ICS+LABA vs. ICS (SMD –0.31, 95% CI –0.48 to –0.13; p = 0.02; 4 trials), and improvement occurred in percentage of symptom-free days (WMD 10.74%, 95% CI 1.86 to 19.62; p=0.02; 3 trials). No difference was found in use of rescue SABAs.	No difference was found between groups in risk of withdrawal, withdrawal due to poor asthma control, risk of AE, and withdrawal due to AE.	

	Stu		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety
agonists versus placebo in addition to inhaled corticosteroids in children and adults	Purpose/Objective: To assess the resulting from the addition of LABAs whether the benefit was influenced buse of 1 or 2 devices to deliver combine the duration of intervention	tients and to examine y obstruction, dose of ICSs, se and type of LABA, and	Addition of LABA improved FEV ₁ (WMD 170 mL, 95% CI 110 to 240, p <0.001).		*Addition of LABA reduced the risk of experiencing ≥1 exacerbations requiring systemic corticosteroids (RR 0.81, 95% CI 0.73 to 0.90, p <0.00005) and increased the	No difference was found in risk of overall AE (RR 0.98, 95% CI 0.92 to 1.05) or withdrawals due to AE (RR 1.29, 95% CI 0.96 to 1.75).	
with chronic asthma. Cochrane Database of Syst Rev 2005; (4):CD0055355. (Canadian Cochrane Network, McGill University, Canada)	Arm 1 LABA + ICS Salmeterol (14 comparisons) or and formoterol (17 comparisons). Arm 2 ICS alone Budesonide (7 trials), beclomethasone (3 trials), budesonide or beclomethasone (1 trial), and uticasone propionate (4 trials); 11 trials failed to specify the ICS.	Most trials used a usual dose of LABA (salmeterol, 50 mcg twice daily; or formoterol, 6 or 12 mcg twice daily). Three trials used 100 mcg twice daily of salmeterol or 24 mcg twice daily of formoterol. One comparison used the 2 options only once daily. Twelve used low-dose ICS (200–400 mcg/day of beclomethasone or equivalent, 8 used a medium dose of ICS (401–799 mcg/day of beclomethasone or equivalent), and 3 comparisons used a high dose of ICS (800 mcg/day of beclomethasone or equivalent).	Nineteen trials used 2 inhaler devices, 5 comparisons used 1 device, 1 tested both 1 and 2 devices, and 1 trial failed to report the number of devices. Duration of intervention was 12–16 weeks (13 trials), 4–8 weeks (6 trials), or 24–54 weeks (7 trials).			proportion of symptom-free days (WMD 17%, 95% CI 12 to 22, p=0.00001; 6 trials) and rescue-free days (WMD 19%, 95% CI 12 to 26, p <0.001; 2 trials). Every 10% increase in baseline FEV ₁ was associated with a 14% increased protection (RR 0.86, 95% CI 0.74 to 1.0) from exacerbations with LABA over placebo.	

	Stu	dy Characteristics		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
Weiler et al. Effect of fluticasone/salmeterol administered via a single device on exercise-induced bronchospasm in patients with persistent asthma. Ann Allergy Asthma Immunol 2005; 94(1):65–72.	Purpose/Objective: To evaluate th fluticasone/salmeterol (F/S) vs. flutic preventing exercise-induced broncher receiving moderate-dose ICSs for the	e effectiveness of regula asone alone (F) adminisospasm in symptomatic	ar treatment with stered via Diskus on adolescents and adults t asthma 4 weeks after 2- to 5 week run-in Albuterol as reliever medication	*At day 1, 1 and 8.5 hr after first dose, maximal decline in FEV ₁ was 11.4% and 11.6%, respectively, for F/S and 20.0% and 12.6%, respectively, for F (p <0.001 and p=0.44). At week 4, 1 and 8.5 hr after last dose, maximal decline in FEV ₁ was 10.9% and 8.9%, respectively, for F (p <0.01 for both). F/S vs. F had greater increase in morning PEF	Values	F/S vs. F had greater increase in percentage of rescue-free days	7% of F/S group and 4% of F group reported AE that was considered drug-related. No SAE was reported.	
				morning PEF (19.2 L/min vs. 6.3 L/min, p=0.03).				

	Stu	dy Characteristics		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
ICS + LTRA vs. ICS								
Robinson et al. Addition of leukotriene antagonists to therapy	Purpose/Objective: To assess who or lung function in patients with chro corticosteroids			No effects for morning or evening PEF or diurnal variation in PEF		No differences in symptom scores or use of rescuer inhaled beta ₂ -agonist use	31% reported AE: 18% while on active treatment and 14% while on placebo.	
in chronic persistent asthma: a randomized double-blind placebo- controlled trial. Lancet 2001;357(9273): 2007–2011.	Arm 1 Montelukast followed by placebo (n=53; 50 completers) Arm 2 Placebo followed by montelukast (n=47; 41 completers)	10 mg montelukast sodium and matched placebo capsules	2 weeks followed by crossover, with no washout period	No difference in FEV ₁ measured at the clinic		There were 4 responders to montelukast and 7 responders to placebo, defined as ≥15% in mean peak flow readings.		
Simons et al. Montelukast added to budesonide in children				*Mean increase in FEV ₁ was 4.6% during montelukast	Blood eosinophil decreased by 15% for	Decrease in beta ₂ -adrenergic agonist use was greater for montelukast vs. placebo (p=0.013).	No difference occurred in incidence of possible drug-related AE (3% montelukast and 3% placebo	
blind, crossover study. J Pediatr 2001; 128:694–698.	Arm 1 Montelukast + budesonide followed by placebo + budesonide Arm 2 Placebo + budesonide followed by montelukast + budesonide (study n=279 randomized; n=264 completers)	5 mg chewable montelukast tablet or matching placebo, 1 daily at bedtime + 200 mcg budesonide twice daily	no washout period During 4-week run-in, received open-label 200 mcg budesonide twice daily	treatment and 3.3% during placebo (diff 1.3%, 95% CI –0.1 to 2.7, p=0.062). Over last 14 days of each period, montelukast vs. placebo difference was 9.7 L/min for morning PEF (95% CI 1.4 to 18.1, p=0.023) and 10.7 L/min for evening PEF (95% CI 2.4 to 19.0, p=0.012).	montelukast group vs. 7% for placebo group (p <0.001).	Percent of asthma exacerbation days was lower during montelukast than placebo treatment (12.2% vs. 15.9%, p <0.001). Montelukast and placebo did not differ in quality of life measurement.	treatment).	

	Stu		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety
Vaquerizo et al. Effect of montelukast added	Purpose/Objective: To evaluate the dose of inhaled budesonide for treat			Morning PEF changed 11.3 in		were lower with montelukast vs.	Eight patients in placebo group and 6 in the montelukast group
to inhaled budesonide on control of mild to moderate asthma. Thorax 2003;58(3): 204–210. (Merck Sharp & Dohme, Spain)	Arm 1 Montelukast (n=313; 308 completers) Arm 2 Placebo (n=326; 317 completers)	10 mg at bedtime	16 weeks after 2-week single-blind placebo run-in Randomization was by stratified budesonide dose level. Salbutamol was used as needed throughout.	placebo and 16.9 in montelukast groups (p=0.05). FEV ₁ increased 2.49% in placebo and 2.63% in montelukast groups (p=0.91).		with relative reduction in risk 21.9% (95%Cl 20.1 to 23.6). Median asthma-free days were greater with montelukast than	discontinued treatment because of clinical AE. Incidence of AE did not differ between groups (40.6% with placebo and 44.2% with montelukast, p=0.37).

	Stu	dy Characteristics		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
ICS + LABA vs. ICS +	LTRA							
Nelson et al. Fluticasone propionate/salmeterol combination provides more effective asthma control than low-dose inhaled corticosteroid plus montelukast. J Allergy Clin Immunol 2000;106(6): 1088–1095. (GlaxoWellcome Inc.)	Purpose/Objective: To compare the propionate/salmeterol combination (montelukast added to fluticasone prosuboptimally controlled with ICS the	ıs inhaler versus	*Overall morning PEF improved in FP/S vs. FP/M (+24.9 L/min vs.		Greater increase occurred in percentage of days with no albuterol use in FP/S group vs. FP/M group (+26.3 vs. +19.1, p=0.032).	AE profiles were similar. Three SAE, occurred, but none were drug-related.		
	Arm 1 Fluticasone propionate + salmeterol + placebo montelukast (FP/S) (n=222; 198 completers) Arm 2 Fluticasone propionate +	100/50 mcg in 1 inhalation twice daily 100 mcg in 1 inhalation twice daily + 10 mg once	in Albuterol was used for	+13.0 L/min, p <0.001). Evening PEF improved in FP/S vs. FP/M (+18.9 L/min vs. +9.6 L/min, p <0.001). FEV ₁ improved in FP/S vs. FP/M (+0.34 vs. +0.20,	Reduction occurred in total albuterol use occurred for FP/S group vs. FP/M group (-1.55 vs1.14 puffs/day, p=0.014).			
	montelukast (FP/M) (n=225; 196 completers)	daily		p <0.001).				
Fish et al. Salmeterol powder provides significantly better benefit than	Purpose/Objective: To compare the LTRA montelukast as add-on therap receiving low-to-intermediate dosage		*S group had greater increase in morning PEF vs. M (35.0 L/min vs. 21.7		S group had greater increase in percentage of symptom-free days vs. M group (24% vs. 16%, p <0.001).	Patients with drug-related AE were 7% in S group and 6% in M group. No drug-related SAE occurred.		
montelukast in asthmatic patients receiving concomitant inhaled corticosteroid therapy. Chest 2001;120(2):423–430. (GlaxoWellcome Inc.)	Arm 1 Salmeterol xinafoate via multidose powder inhaler + ICS (S) (n=476) Arm 2 Montelukast + ICS (M) (n=472)	50 mcg via multidose powder inhaler + usual dose ICS 10 mg/day orally + usual dose ICS	12 weeks after 7-day to 14-day run-in period Albuterol inhaler for relief of breakthrough symptoms	L/min, p <0.001). Improvements were noted within 1st week and remained over 12 treatment weeks.		Daytime symptom scores decreased by 39% in S group vs. 31% in M group (p=0.039). Supplemental albuterol use was less in S group vs. M group (42% vs. 35% for daytime; 51% vs. 40% for nighttime; p <0.012). Greater reduction in nighttime awakenings/week occurred in S group vs. M group (1.42 vs. 1.32, p=0.015).		

	Stu	dy Characteristics		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
Nelson et al. Comparison of inhaled salmeterol and oral zafirlukast in asthmatic patients using concomitant inhaled corticosteroids. MedGenMed	Purpose/Objective: To compare the addition of zafirlukast to a regimen of Arm 1 Salmeterol xinafoate (S)	e effects of the addition f ICSs 42 mcg (2 puffs) twice daily via metered dose inhaler + stable dose of ICS 20 mg twice daily +	of salmeterol vs. the 4 weeks after 7- to 14-day run-in period	*S produced greater increase in mean morning PEF (28.8 L/min) and evening PEF (21.8 L/min) vs. Z (13.0 L/min and 11.2 L/min, p <0.001 and p=0.004, respectively). Decrease in mean PEF differential was greater for S vs. Z (-8.1 L/min vs3.7 L/min, p=0.022). Greater improvement in PEF occurred for S group		Daytime symptom scores decreased by 35% in S group vs. 21% in Z group (p=0.002). S group vs. Z group had greater	Percentage reporting ≥1 AE in each group was 39%. In each group, 7 patients (3.3%) withdrew due to AE; 5 AE in the Z group were study-drug related.	
				vs. Z group at all treatment weeks. Increase in mean predose FEV ₁ was greater for S group vs. Z group at week 1 (0.23 vs. 0.16, p <0.05); no difference occurred at week 4.		Improvement in AQLQ was greater in S group vs. Z group (p <0.009). In each group, 8 patients (3.7%) experienced exacerbations.		

	Stu	Study Characteristics			Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety		
Bjermer et al. Montelukast and fluticasone compared with salmeterol and fluticasone in protecting against asthma exacerbation in adults: one year, double blind, randomised, comparative trial. BMJ 2003;327(7420):891. (Merck & Co.)	Purpose/Objective: To assess the inhaled fluticasone propionate on as inadequately controlled with fluticasonal Arm 1 Montekulast (M/F) (n=747; 83.3% completed) Arm 2 Salmeterol (S/F) (n=743; 85.2% completed)	thma exacerbation in pa	48 weeks after 4-week run- in period	No difference between groups in change in FEV ₁ (0.11 vs. 0.19). M/F group showed smaller decrease in percent reversibility in FEV ₁ (-7.54 vs11.26, p <0.001). S/F group showed larger increase in PEF (34.59 vs. 17.73, p <0.001).	M/F treatment reduced peripheral blood eosinophil counts from baseline (-0.04 10³/mcL, p <0.001) but S/F treatment did not (-0.01, p >0.05).		AE were reported by 71.0% of M/F group and 72.4% of S/F group. S/F group had a higher incidence of drug-related AE (7.4% vs. 4.6%, p=0.022).		

	Stu	dy Characteristics		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
Ringdal et al. The salmeterol/fluticasone combination is more effective than fluticasone plus oral montelukast in asthma. Respir Med 2003;97(3):234–241. (GlaxoSmithKline)	Purpose/Objective: To compare the or montelukast to treatment with an I who are symptomatic despite treatment. Arm 1 Fluticasone propionate + oral montelukast (FP+M) (n=369 in ITT analysis) Arm 2	e clinical effect of the ad CS, fluticasone propion	ddition of either salmeterol ate, in adults with asthma 12 weeks treatment + 2-week followup after 4-week run-in period Salbutamol was used for rescue relief as required. Other regular asthma medication continued at constant dose.	*Adjusted mean increase in morning PEF was greater in SF group vs. FP+M group over 12 weeks (36 L/min vs. 19 L/min; diff. 17 L/min, 95% CI 12 to 22 L/min, p <0.05). Onset of improvement was faster in SF group, with difference at 24 hours (diff 16.9 L/min, 95% CI 11.9 to 32 L/min, p=0.03). Increase in FEV ₁ was greater for SF group vs. FP+M group (0.26 vs. 0.17; diff. 0.11 L, 95% CI 0.06 to 0.16 L.	values	SF group was more likely to have symptom-free day during study period (OR 1.32, 95% CI 1.05 to 1.65, p <0.04) and more likely to have rescue-free day (OR 1.39, 95% CI 1.02 to 1.64, p=0.03). 9.6% of SF group vs. 14.6% of FP+M group had at least 1 exacerbation (p <0.05). Time to 1st exacerbation was longer in SF group than in FP+M group (p <0.05).	Similar incidence of AE occurred (44% of SF group vs. 42% of FP+M group). No drug-related SAE occurred.	

	Stu		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety
Ceylan et al. Addition of formoterol or montelukast to low-	Purpose/Objective: To investigate a low dose of ICS was effective in the cases and to determine which drug s	e control of asthma in m		(Analysis was adjusted for gender, age, and baseline		(Analysis was adjusted for gender, age, and baseline values.) Decrease in beta ₂ -agonist use was	Local AE effects potentially related to drugs occurred for 30% of FB group and 20% of MB group.
dose budesonide: an efficacy comparison in short- and long-term asthma control. Respiration 2004;71(6):594–601.	Arm 1 Formoterol + budesonide (FB) (n=20 completers) Arm 2 Montelukast + budesonide (MB) (n=20 completers)	9 mcg twice daily + 400 mcg 10 mcg once daily + 400 mcg	Salbutamol, 100 mcg/puff, was allowed for treatment of symptoms.	values.) *Morning PEF increased from 266.3 to 320.5 L/min in FB group vs. 262.8 to 293.3 L/min in MB group (diff. of 23.7 L/min increase, p <0.0001). Night PEF increased from 287 to 331.5 L/min in FB group and from 283 to 310 L/min in MB group (diff of 17.5 L/min increase, p <0.001). Improvements in FB group began earlier. FEV ₁ change was greater in FB vs. MB group (0.36 vs. 0.19, p <0.001). FEV ₁ % pred. change was greater in FB vs. MB group (10.7 vs. 5.6, p <0.001).		greater in FB group vs. MB group (1.9 vs. 0.5 puffs/day, p <0.001). Decrease in morning symptom scores was greater for FB group vs. MB group (2.6 vs. 0.8, p <0.0001).	

	Stu	dy Characteristics		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
montelukast or salmeterol to	Purpose/Objective: To compare madministered with inhaled fluticason asthma attack for 1 year			reduced %	Montelukast vs. salmeterol reduced blood	*20% of M/F group and 16.7% of S/F group had asthma attacks (RR=1.20, 95% CI 0.96 to 1.49).	Monetlukast and salmeterol groups were comparable in proportion with clinical AE that was possibly or	
fluticasone for protection against asthma attacks: a randomized, doubleblind, multicenter study. Ann Allergy Asthma Immunol 2004;92(6):641–648.	Arm 1 Montelukast + fluticasone MDI (M/F) (n=743; 734 in analysis; 530 completers)	10 mg/day + 220 mcg/day fluticasone	·	reversibility in FEV ₁ (diff. = 4.78 , 95% CI 3.89 to 5.67). Salmeterol vs. montelukast treatment increased FEV ₁ (diff. 1.98,			definitely drug related (8.6% vs. 10.0%) and serious (3.0% vs. 3.7%).	
	Arm 2 Salmeterol MDI + fluticasone MDI (S/F) (n=730; 718 in analysis; 529 completers)	42 mcg twice daily + 220 mcg/day fluticasone		95% CI 1.01 to 2.96) and PEF (diff. 14.3, 95% CI 6.4 to 22.1).				
Ram et al. Long-acting beta ₂ -agonist versus	Purpose/Objective: To compare the in asthmatic patients who remain sys			ICS+LABA vs. ICS+LTRA group		*Risk of exacerbation requiring systemic corticosteroids was lower	Risk of withdrawal was reduced for ICS+LABA vs. ICS+LTRA (RR 0.84,	
anti-leukotrienes as add-on therapy to inhaled corticosteroids for chronic asthma. Cochrane Database of Syst Rev 2005;(1):CD003137. (NHS Research and Development UK) NOTE: Meta-analysis includes studies abstracted separately above: Bjermer et al. 2003; Fish et al. 2001; llowite et al. 2004; Nelson et al. 2000; and Nelson et al.	Arm 1 ICS+LABA Arm 2 ICS+LTRA		(1 trial), 12 weeks (4 trials), and 48 weeks (2 trials).	showed greater improvement in morning PEF (WMD 15.75 L/min, 95% CI 13.0 to 18.5; 8 trials) and in evening PEF (WMD 11.86 L/min, 95% CI 8.85 to 14.86; 7 trials). Improvement in FEV ₁ favored ICS+LABA group (WMD 0.08 L, 95% CI 0.06 to 0.10; 7 trials).			95% CI 0.74 to 0.96; 8 trials), with no difference in withdrawals due to AE (RR 1.03, 95% CI 0.80 to 1.33; 8 trials).	

	Stu	dy Characteristics		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
ICS + LTRA vs. increa	sing ICS							
Price et al. A randomized controlled trial of montelukast	Purpose/Objective: To compare the budesonide with doubling the dose of symptomatic on inhaled budesonide		*Improvement in morning PEF over last 10 weeks with M/B was as effective	Groups did not differ in change in peripheral blood	nocturnal awakenings (p=0.353),	No difference was found between groups in number with AE, drug-related AE, SAE, or discontinuing because of AE.		
plus inhaled budesonide versus double dose inhaled budesonide in adult patients with asthma.	Arm 1 Montelukast + budesonide (M/B) (n=448; 428 completers)	Montelukast, 10 mg/day at bedtime + budesonide, 800 mcg/day	12 weeks after 4-week run- in period	as B (33.5 L/min vs. 30.1 L/min, 95% CI –12.9 to 4.8 for difference).		median days with exacerbations (6.7% vs. 6.3%, p=0.78), median asthma-free days (86.7% vs. 82.2%, p=0.37), or proportion requiring oral steroids or admission		
Thorax 2003;58(3):211–216. (Merck & Co.)	Arm 2 Budesonide (B) (n=441; 415 completers)	Budesonide, 800 mcg/day, twice daily	SABAs were used on as- needed basis.	Change during first 3 days in morning PEF was greater in M/B group than in B group (20.1L/min vs. 9.6 L/min; 95% CI –17.6 to –4.3, p <0.001).		to hospital (1.6% vs. 2.3%, p=0.47).		

Stu	Findings					
Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety
ising ICS						
Purpose/Objective: To compare the fluticasone propionate combination of patients with moderate-to-severe permoderate-to-high corticosteroid doses Arm 1 Salmeterol + fluticasone propionate (SFC) (n=180, 151 completers)	vith a threefold higher manager is stent asthma who rendered by the state of the st	24 weeks after 2-week run- in Rescue salbutamol was used as needed throughout.	morning PEF (adjusted for age, gender, country, and baseline) was found for SFC vs. B treatment (406 L/min vs. 380 L/min, diff. 25 L/min, 95% CI 15 to 35, p <0.001). Greater improvement occurred in adjusted evening PEF for SFC vs. B group (416 L/min vs. 398 L/min, p <0.001). Adjusted mean diurnal variation in PEF was lower in SFC vs. B group (p <0.003) but was		Percentage of symptom-free days over 24 weeks was greater in SFC group vs. B group (95% CI 2 to 11, p <0.001). SFC group had greater reduction than B group in rescue medication use (p <0.001).	14% of SFC group and 18% of B group reported treatment-related AE. No treatment-related SAE occurred.
	Treatment Psing ICS Purpose/Objective: To compare the fluticasone propionate combination of patients with moderate-to-severe permoderate-to-high corticosteroid doses Arm 1 Salmeterol + fluticasone propionate (SFC) (n=180, 151 completers) Arm 2 Budesonide (B)	Purpose/Objective: To compare the efficacy and tolerability fluticasone propionate combination with a threefold higher may patients with moderate-to-severe persistent asthma who remoderate-to-high corticosteroid dose Arm 1 Salmeterol + fluticasone propionate (SFC) (n=180, 151 completers) Arm 2 Budesonide (B) 800 mcg plus placebo twice daily	Treatment Dose Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup sing ICS Purpose/Objective: To compare the efficacy and tolerability of this salmeterol plus fluticasone propionate combination with a threefold higher microgram dose of ICS in patients with moderate-to-severe persistent asthma who remain symptomatic on a moderate-to-high corticosteroid dose Arm 1 Salmeterol + fluticasone propionate (SFC) (n=180, 151 completers) Arm 2 Budesonide (B) (n=173; 143 completers) Budesonide (B) (n=173; 143 completers)	Treatment Dose Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup Sing ICS Purpose/Objective: To compare the efficacy and tolerability of this salmeterol plus fluticasone propionate combination with a threefold higher microgram dose of ICS in patients with moderate-to-severe persistent asthma who remain symptomatic on a moderate-to-high corticosteroid dose Arm 1 Salmeterol + fluticasone propionate (SFC) (n=180, 151 completers) Arm 2 Budesonide (B) (n=173; 143 completers) 800 mcg plus placebo twice daily 800 mcg plus placebo twice daily 800 mcg plus placebo twice daily Creater improvement occurred in adjusted evening PEF for SFC vs. B group (416 L/min vs. 398 L/min, p <0.001). Adjusted mean diurnal variation in PEF was lower in SFC vs. B group	Treatment Dose Duration of Active Treatment; Dose Duration of Postintervention/ Off-Treatment Followup Lung Function Lung Function Lung Function Vital Signs/ Cardiovascular/ Clinical Laboratory Values **Significant difference in morning PEF (adjusted for age, gender, country, and baseline) was found for SFC vs. B treatment (406 L/min vs. 380 L/min, 95% Cl 15 to 35, p < 0.001). Arm 2 Budesonide (B) (n=173; 143 completers) **Significant difference in morning PEF (adjusted for age, gender, country, and baseline) was found for SFC vs. B group (416 L/min vs. 398 L/min, p < 0.001). **Greater improvement occurred in adjusted evening PEF for sproup (416 L/min vs. 398 L/min, p < 0.001). Adjusted mean diurnal variation in PEF was lower in SFC vs. B group (p < 0.003) but was not clinically was not cli	Treatment Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup Sing ICS Purpose/Objective: To compare the efficacy and tolerability of this salmeterol plus fluticasone propionate combination with a threefold higher microgram dose of ICS in patients with moderate-to-severe persistent asthma who remain symptomatic on a moderate-to-fluticate for age, gender, country, and basel, patients with moderate-to-fluticasone propionate (SFC) (n=180, 151 completers) Arm 2

	Stu	dy Characteristics		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
salmeterol to low-dose fluticasone versus higher-dose	Purpose/Objective: To compare the in patients after adding salmeterol to rates and characteristics of exacerba propionate (F)	low-dose fluticasone pi	ropionate (S/F) with the ng higher dose fluticasone	Changes in morning PEF during exacerbation were comparable in the		*8.8% of S/F group vs. 13.8% of F group had ≥1 exacerbation (p=0.017). Characteristics of those with		
exacerbations. J Allergy Clin Immunol 2001;107(5):783–789.	Arm 1 Salmeterol + fluticasone (S/F) (n=467) Arm 2 Fluticasone (F) (n=458)	42 mcg/88 mcg twice daily 220 mcg twice daily	24 weeks, after 2- to 4-week screening period Albuterol was used to relieve break-through symptoms.	2 groups.		exacerbations were similar in the 2 groups. Mean duration of exacerbation was 8.4 days in S/F group and 10.5 days in F group (p=0.17) and required 6.6 days vs. 7.5 days of treatment (p=0.12). Time to 1st exacerbation favored S/F group (p=0.049). Rescue albuterol use did not differ between the 2 groups.		

	Stu	Findings					
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety
O'Byrne et al. Low dose inhaled budesonide and	Purpose/Objective: To determine budesonide with or without low dose exacerbations and improve asthma	Group A (adjusted for baseline) BF vs. B treatment		Group A (adjusted for baseline) *B vs. P group showed reduction in risk for 1st severe exacerbation	Number of AE was similar between different treatments.		
formoterol in mild persistent asthma: the OPTIMA randomized trial. Am J Respir Crit Care Med 2001;164(8 Pt 1):1392–1397.	Group A Arm 1 Budesonide (B) (n=228) Arm 2 Budesonide + formoterol (BF) (n=231) Arm 3 Placebo (P) (n=239) Group B Arm 1 Budesonide (100B) (n=228) Arm 2 Budesonide + formoterol (100BF) (n=231) Arm 3 Budesonide (200B) (n=239) Arm 4 Budesonide + formoterol (200BF)	100 mcg twice daily 100/4.5 mcg twice daily placebo 100 mcg twice daily 100/4.5 mcg twice daily 200 mcg twice daily	Group B took budesonide, 100 mcg twice daily) No additional treatments were allowed. If a patient had a severe exacerbation, medications were at the physician's discretion.	increased FEV ₁ % pred. (5.87 vs. 4.04, p <0.005) and resulted in greater change in morning PEF (31.81 vs. 15.12, p <0.001). Group B (adjusted for baseline) Adding formoterol increased FEV ₁ and morning PEF (p <0.001). 100BF vs. 200B treatment improved FEV ₁ (p=0.05) and morning PEF (p <0.005).		(RR=0.40, 95% CI 0.27 to 0.59). B vs. P group reduced rate of poorly controlled asthma days (RR=0.52, 95% CI 0.40 to 0.67), rate of exacerbations (RR=0.38, 95% CI 0.25 to 0.57), days with asthma symptoms (p <0.001), days with nocturnal awakening (p <0.001), and number of rescue inhalations (p <0.001). Group B (adjusted for baseline) *200B vs. 100B group had reduced risk of 1st severe exacerbation (RR=0.81, 95% CI 0.65 to 1.01). 200B vs. 100B reduced rate of poorly controlled asthma days (RR=0.87, 95% CI 0.75 to 1.01). Adding F reduced risk of 1st exacerbation (RR=0.57, 95% CI 0.46 to 0.72) and rate of severe exacerbations (RR=0.48, 95% CI 0.69 to 0.59). 100BF was more effective than 200B in reducing risk of severe exacerbation day (RR=0.71, 95% CI 0.52 to 0.96) and rate of severe exacerbations (RR=0.58, 95% CI 0.44 to 0.76).	

	Stu		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety
Bateman et al. Combination therapy with single inhaler budesonide/formoterol	Purpose/Objective: To compare the twice daily with a high dose of the condaily in patients with moderate persistence.	orticosteroid fluticasone	propionate 250 mcg twice	*Greater increase in morning PEF occurred with B/F vs. FP (27.4 vs.		lower with B/F vs. FP (diff 0.18, 95% CI 0.01 to 0.35, p=0.04).	Similar AE profiles for occurred with B/F and FP. No treatment-related SAE occurred.
compared with high dose of fluticasone propionate alone in patients with moderate persistent asthma. Am	Arm 1 Budesonide/formoterol (B/F) (n=168; 153 completers) Arm 2 Fluticasone propionate (FP) (n=176; 156 completers)	160/4.5 mcg twice daily 250 mcg twice daily	12 weeks, after 2-week run-in period Terbutaline sulfate or albuterol was used as reliever medication.	7.7 L/min, p <0.001). Difference was evident on 1st day (p <0.001) and continued over last 30 days (p <0.001).		Reliever-free days were increased with B/F vs. FP (75.5 vs. 66.4, diff. 9.1, 95%Cl 3.8 to 14.3, p <0.001). No difference was found in nighttime awakenings, symptom-free days, and asthma-control days. Risk of mild exacerbation was 32% lower in B/F (RR 0.678, 95% Cl 0.465 to 0.988).	
Ind et al. Addition of salmeterol to fluticasone propionate	Purpose/Objective: To see whether that of doubling the dose of fluticaso who continued treatment with low-do	ne propionate while also	including a control group	*Improvement in PEF was greater with S/F (42 L/min)		*66% of S/F, 59% of F250 and 65% of F500 groups had exacerbations (p >0.05).	
Pospir Mod	(S/F) (n=171; 144 completers)	Salmeterol 50 mcg twice daily + fluticasone 250 mcg twice daily Fluticasone 250 mcg twice daily Fluticasone 500 mcg twice daily	24 weeks, after 4-week run-in period Salbutamol was used for symptom relief, and oral prednisolone was used in exacerbations.	compared with F500 (16.5 L/min) and F250 (16.9 L/min), p <0.001. No difference was found between F250 and F500. S/F reduced diurnal variation in PEF more (-4.9%) vs. F500 (-3.0%) and F250 (-2.2%), both p <0.001.		No difference was found in percentage of patients experiencing moderate or severe exacerbations (28% of S/F, 31% of F500, and 25% of F250). Proportion of symptom-free days and nights increased more with S/F (median 21%) compared with F500 (median 1.5%) and F250 (median 0%), both p=0.002.	

Duration of Active Treatment; Cardiovascular/ Duration of Clinical	
Citation/ Sponsor Treatment Dose Postintervention/ Off-Treatment Followup Lung Function Values Symptoms	Safety
Lalloo et al. Budesonide and formoterol in a single inhaler improves Purpose/Objective: To evaluate the efficacy and safety of low-dose budesonide (B) in adult formoterol in a single inhaler improves Purpose/Objective: To evaluate the efficacy and safety of low-dose budesonide (B) in adult formoterol (B/F) compared with increased dose of budesonide (B) in adult formoterol in a single inhaler improves Purpose/Objective: To evaluate the efficacy and safety of low-dose budesonide (B) in adult formoterol (B/F) compared with increased dose of budesonide (B) in adult formoterol in a single inhaler improves Purpose/Objective: To evaluate the efficacy and safety of low-dose budesonide (B) in adult formoterol (B/F) compared with increased dose of budesonide (B) in adult formoterol (B/F) compared with increased by 17% in B/F group and 10% in B group (p=0.002). Proportion of asthma-control days increased by 17% in B/F group and 10% in B group (p=0.002). Proportion of symptom-free days Proportion of sym	nce was found between frequency of AE; 5 SAE red with B/F treatment to treatment) and 2 SAE with B treatment.

Study Characteristics				Findings			
Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
			*SF was superior to F treatment in morning PEF (48 L/min vs.	Blood pressure and heart rate remained stable throughout	SF group vs. F group had an increased percentage of symptom-free days (+40 vs. +29, adj. diff. 12.8, 95% CI 4.5 to 21.0, p=0.0025	26.3% AE occurred in SF group vs. 24.2% in F group. One possible drug-related SAE	
meterol + fluticasone (SF) 179; 166 completers)	daily	12 weeks, after 2-week screening period Salbutamol in metered dose inhalers was used as rescue medication.	30 L/min, diff. 19.6, 95% CI 6.8 to 32.4, p=0.0027 at 6 weeks; +52 L/min vs. +36 L/min, diff. 16.6, 95% CI 1.1 to 32.0, p=0.036 at 12 weeks). Difference was adjusted for baseline, age, sex, height, and duration of preceding treatment with ICS. No difference	treatment.	12.8, 95% CI 4.5 to 21.0, p=0.0025 at 6 weeks; +49 vs. +38, adj. diff. 12.6, 95% CI 4.0 to 20.7, p=0.004 at 12 weeks). Rescue medication use decreased more in SF group vs. F group (-1.4 vs1.0, adj. diff -0.5, 95% CI -0.85 to -0.20, p=0.0015 at 6 weeks; -1.6 vs1.0, adj. diff0.84, 95% CI -1.13 to -0.37, p <0.001 at 12 weeks.	occurred in F group vs. 2 unrelated SAE in SF group.	
nb de n 17	Treatment ose/Objective: To study the efination (SF) in comparison witherate symptomatic asthma 1 eterol + fluticasone (SF) 79; 166 completers) 2 asone(F)	Treatment Dose ose/Objective: To study the efficacy and tolerability orbination (SF) in comparison with doubling the dose of fluorate symptomatic asthma 1 eterol + fluticasone (SF) 79; 166 completers) 2 asone(F)	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup ose/Objective: To study the efficacy and tolerability of the salmeterol fluticasone ination (SF) in comparison with doubling the dose of fluticasone (F) in patients with erate symptomatic asthma 1	Treatment Dose Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup See/Objective: To study the efficacy and tolerability of the salmeterol fluticasone ination (SF) in comparison with doubling the dose of fluticasone (F) in patients with parate symptomatic asthma 1	Treatment Dose Direction of Active Treatment; Duration of Postintervention/ Off-Treatment Followup Dose/Objective: To study the efficacy and tolerability of the salmeterol fluticasone ination (SF) in comparison with doubling the dose of fluticasone (F) in patients with parate symptomatic asthma 1	Treatment Dose D	

	Study Characteristics			Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
Jonsson et al. An economic evaluation of combination treatment with budesonide and formoterol in patients with mild-to-moderate persistent asthma. Respir Med 2004; 98(11):1146–1154.	Purpose/Objective: To present the results of an economic analysis of 3 step-up treatments					B100/F was more expensive and less effective than B200/F and could not provide best "value for		
	Arm 1 Budesonide (B100) (n=322) Arm 2 Budesonide (B200) (n=312)	100 mcg twice daily 200 mcg twice daily	1 year, after 4-week run-in			money." B200/F provided more symptom- free days but was associated with higher costs than B200. The incremental cost-effectiveness ratio for this comparison was Sweden Kronor 21 per symptom-free day gained.		
	Arm 3 Budesonide/formoterol (B100/F) (n=323)	100/4.5 mcg twice daily						
	Arm 4 Budesonide/formoterol (B200/F) (n=315)	200/4.5 mcg twice daily						

	Stu	dy Characteristics		Findings				
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety	
Masoli et al. Moderate dose inhaled corticosteroids plus	Purpose/Objective: To compare the controlled on moderate doses of ICS			Those who received low-dose ICS/salmeterol vs.			*Decreased number of subjects withdrew due to asthma in low-dose ICS/ salmeterol vs. high-dose ICS	
salmeterol versus higher doses of inhaled	Salmeterol + BDP 400 mcg/day vs. BDP 1,000 mcg/day (1 study)		12 weeks (7 studies), 24 weeks (5 studies), or 26	high-dose ICS had reduced morning			treatment (2.9% vs. 4.3%; OR 1.58, 95% CI 1.12 to 2.24).	
	Salmeterol + BDP 400 mcg/day vs. BDP 800 mcg/day (3 studies)			PEF (diff. 23 L/min, 95% CI 10 to 28) and evening PEF			*Reduced number of subjects had ≥1 moderate or severe exacerbation	
2005;60(9):730–734.	Salmeterol + fluticasone 200 mcg/day vs. fluticasone 500 mcg/day (6 studies)				(diff. 19 L/min, 95% CI 15 to 23), greater daytime			in low-dose ICS/salmeterol vs. high-dose ICS treatment (8.0% vs. 10.7%; OR 1.35, 95%Cl 1.10 to 1.66).
	Salmeterol + fluticasone 200 mcg/day vs. budesonide 800 mcg/day (1 study)			beta ₂ -agonist use (diff. –0.58, 95% CI –0.02 to –0.13), and			1.00).	
				reduced FEV ₁ (diff. 0.12 L, 95% CI 0.09 to 0.15).				

	Study Characteristics			Findings			
Citation/ Sponsor	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Lung Function	Vital Signs/ Cardiovascular/ Clinical Laboratory Values	Exacerbations/ Symptoms	Safety
LABA A vs. LABA B in a	addition to ICS						
of bronchodilation of a	Purpose/Objective: To evaluate the and formoterol in a single inhaler in of fluticasone	comparison with the con	nbination of salmeterol and	*Both BF groups had faster onset of improvement in FEV ₁ compared to			
fluticasone in single inhalers. Pulm Pharmacol Ther 2001;14(1):29–34.	Arm 2 Budesonide/formoterol (BF2)	160/4.5 mcg (2 inhalations) 50/250 mcg	least 8 hours, LABAs were withheld at least 72 hours, and LTRAs and anticholinergics were withheld at least 12 hours prior to study drug.	FEV ₁ compared to SF at 3 minutes after dose (2.74 and 2.75 vs. 2.56, p <0.001) and at 0-5 minute average FEV ₁ (2.80 and 2.83 vs. 2.67, p <0.001). No evidence was found of difference between 2 BF doses for any changes in FEV ₁ up to 3 hours after inhalation. 47% of SF group showed onset of effect (15%) after inhalation of SF within 60 minutes			

	Stu	dy Characteristics		Findings				
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Everden et al. Eformoterol Turbohaler	Purpose/Objective: To examine the compared with salmeterol in children		afety of eformoterol	Both groups showed improvements over		Reductions for E vs. S group were	AE was reported by 55% of E group and 59% of S group. Number,	
compared with salmeterol by dry powder inhaler in asthmatic children not controlled on inhaled corticosteroids. Pediatr Allergy Immunol 2004; 15(1):40–47.	Arm 1 Eformoterol Turbohaler® (E) (n=80; n=79 in analysis; n=59 completers) Arm 2 Salmeterol Accuhaler® (S) (n=76; n=64 completers)	12 mcg twice daily (9 mcg delivered dose) 50 mcg twice daily	12 weeks, after 7- to 10-day run-in period Patients continued to receive current ICS and SABA throughout study.	time in clinic PEF at 4, 8, and 12 weeks (p <0.01).		daytime use and 66% vs. 49% for	nature, and intensity of AEs were the same for both groups. No treatment-related SAE occurred.	