July 2007

Evidence Table 13. Pharmacologic Therapy: Immunomodulators—Anti-Immunoglobulin E

Abbreviations used in table:

AE	adverse event	NNT	num
BDP	beclomethasone dipropionate	Ο	oma
Der f	Dermatophagoides farinae	Р	place
Der p	Dermatophagoides pteronyssinus	$PD_{20}FEV_1$	cum
EAACI	European Academy of Allergology and Clinical Immunology		FEV ₁
FEV ₁	forced expiratory volume in 1 sec	PEF	peak
HD	high dose	QoL	quali
ICS	inhaled corticosteroid	SAE	seve
lgE	immunoglobulin E	SD	stan
ITT	intent-to-treat	SIT	spec
LABA	long-acting beta-agonist		

* indicates primary outcome

NNT	number needed to treat
0	omalizumab
Р	placebo
PD ₂₀ FEV ₁	cumulative dose of methacholine producing a 20% decrease in \ensuremath{FEV}_1
PEF	peak expiratory flow
QoL	quality of life
SAE	severe adverse event
SD	standard deviation
SIT	specific immunotherapy

Evidence Table 13. Pharmacologic Therapy: Immunomodulators—Anti-Immunoglobulin E

		Study Population					
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)			
Boulet et al. Inhibitory effects of an anti-IgE antibody E25 on allergen-induced early asthmatic response. Am J Respir Crit Care Med 1997;155(6):1835–1840. (Genentech Inc.)	Multicenter, randomized, double-blind, parallel-group, placebo-controlled trial	20 (19—1 withdrawal from treatment group)	Age 21-44 yr; mean = 27.4 yr (SD=8.1) Gender 12 male, 8 female Ethnicity Not reported Highly positive allergy skin-prick test to at least 1 common aeroallergen and early asthmatic response on allergen inhalation in lab	Stable, mild, allergic asthma requiring only an inhaled beta ₂ -agonist on demand FEV ₁ % pred. mean = 91.9 (SD=11.07)			
Fahy et al. The effect of an anti-IgE monoclonal antibody on the early- and late- phase responses to allergen inhalation in asthmatic subjects. Am J Respir Crit Care Med 1997;155(6):1828–1834. (Genentech Inc.)	Double-blind, placebo-controlled, randomized parallel group study	19 (18)	Age Mean = 31.5 yr Total serum IgE, mean = 142 IU/L	Mild asthma requiring only inhaled beta ₂ -agonists FEV ₁ % pred., mean = 94			
Milgrom et al. Treatment of allergic asthma with monoclonal anti-IgE antibody. N Engl J Med 1999;341(26):1966–1973. (Genentech Inc.)	Multicenter, randomized, double-blind, placebo-controlled trial	317 (306)	Age 11–50 yr, mean = 30 yr (54 adolescents) Gender 42% male, 58% female Ethnicity Not reported Total serum IgE 17–1,957 IU/mL	Moderate-to-severe perennial allergic asthma $FEV_1 \%$ pred., mean = 71; range, 29–129 Mean asthma symptom score = 4.0 (range, 1.5–6.5 out of 1–7) Daily use of beta ₂ -agonist bronchodilator as rescue medication Median duration of asthma, 19 yr			

		Study Population					
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)			
Busse et al. Omalizumab, anti-IgE recombinant humanized monoclonal antibody, for the treatment of severe allergic asthma. J Allergy Clin Immunol 2001;108(2):184–190. (Novartis Pharmaceuticals Corp. and Genentech, Inc.)	Multicenter, double-blind, placebo- controlled, randomized Phase III trial	525 (525; 53 dropouts)	Age 12–74 yr; mean = 39.2 Gender 41% male, 59% female Ethnicity 89% Caucasian, 7% Black, 4% other Total serum IgE, 20–860 IU/mL (mean = 179)	Severe allergic asthma requiring daily ICS Duration 1–61 yr, mean = 21.6 yr FEV ₁ % pred., mean = 68.0 BDP dose, 336–1008 mcg/day; mean = 569 mcg/day			
Milgrom et al. Treatment of childhood asthma with anti-immunoglobulin E antibody (omalizumab). Pediatrics 2001;108(2):E36. (Genentech, Inc.; Novartis Pharmaceuticals Corp.)	Randomized, double-blind, placebo- controlled, parallel-group design	334 (334)	Age 5–12 yr, mean = 9.4 yr Gender 69.2% male, 30.8% female Ethnicity White, 76.0%, Black, 15.6%, other, 8.4%	Moderate-to-severe allergic asthma Duration of allergic asthma: range, 1–12 yr; mean = 6.1 yr BDP dose: range, 168–672 mcg/day; mean = 278 mcg/day Serum total IgE: range, 20–1269 IU/mL; mean = 340 IU/mL FEV ₁ % pred.: range, 43–129; mean = 84 FEV ₁ reversibility, mean = 20.1% Morning PEF: range, 101–408 L/min; mean = 262 L/min Hospitalized for asthma in past year, 8.1% All children had minimal asthma symptoms, with mean rescue albuterol use <2 puffs/day.			
Soler et al. The anti-IgE antibody omalizumab reduces exacerbations and steroid requirement in allergic asthmatics. Eur Respir J 2001;18(2):254–261. (Novartis Pharma AG and Genentech Inc.)	Multicenter, randomized, double-blind, placebo-controlled, parallel-group study	546 (487)	Age 12–76 yr, mean = 39.5 yr Gender 49% male, 51% female Ethnicity Not reported Total serum IgE: range, 21–814 IU/mL, mean = 214.4 IU/mL	Moderate-to-severe allergic asthma, symptomatic despite ICS (500–1,200 mcg/day BDP) Mean duration of asthma, 19.7 yr; range, 1–68 yr			

			Study Population	
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)
Walker et al. Anti-IgE for chronic asthma. Cochrane Database Syst Rev 2002;(3): CD003559.	Meta-analysis of randomized, double-blind, parallel-group controlled trial	8 trials; n=2,037	Age 6–12 yr (1 trial, n=334), ≥12 yr (7 trials, n=1,703)	Mild (3 trials), moderate to severe (4 trials), and severe asthma (1 trial)
(Garfield Weston Foundation UK ; NHS Research and Development UK ; The Thriplow Charitable Trust UK)	Methodological quality as assessed by Jadad scale was fair to high (three scored 3, three scored 4, two scored 5).			
Berger et al. Evaluation of long-term safety of the anti-IgE antibody, omalizumab, in children with allergic asthma. Ann Allergy Asthma Immunol 2003;91(2):182–188. (Novartis Pharmaceuticals Corp.)	Randomized, double-blind, placebo-controlled trial (only treatment group reported here)	225 (225)	Age 5–12 yr; mean = 9.4 Gender 70% male, 30% female Ethnicity 75% Caucasian, 17% Black, 8% other Total serum IgE: range, 30–1,300 IU/mL (mean = 348); body weight ≤90 kg 18% hospitalized for asthma in past year	204 moderate (FEV ₁ % pred. >65%); 21 severe (FEV ₁ % pred. \leq 65%) Duration: range, 1–12 yr; mean = 6.1 yr Asthma well controlled \geq 3 months with ICS doses equivalent to 168–420 mcg/day of BDP
Corren et al. Omalizumab, a recombinant humanized anti-IgE antibody, reduces asthma-related emergency room visits and hospitalizations in patients with allergic asthma. J Allergy Clin Immunol 2003; 111(1):87–90. (Novartis Pharma AG; Genentech, Inc.)	Pooled analysis of 3 multicenter, randomized, double-blind, placebo- controlled trials	1,405 (1,405)	Age ≥12 yr (2 studies, n=1,071); ages 6–12 yr (1 study, n=334) Gender Not reported Ethnicity Not reported	Moderate-to-severe allergic asthma Duration of asthma, ≥1 yr Total serum IgE level: range, 30–700 IU/mL (adolescents/adults) or 30–1,300 IU/mL (children) Required daily ICSs Positive skin-prick test to dust mite, cockroach, dog, or cat
Ayres et al. Efficacy and tolerability of anti- immunoglobulin E therapy with omalizumab in patients with poorly controlled (moderate-to- severe) allergic asthma. Allergy 2004;59(7): 701–708.	Multicenter, randomized, open-label, parallel-group study 49 centers in 5 European countries	312 (ITT analyses)	Age 12–73 yr, median 38 yr Gender 29% male, 71% female Ethnicity Not reported Serum IgE: range, 27–686 IU/mL; median, 167 IU/mL (treatment group only)	Poorly controlled, moderate-to-severe allergic asthma; receiving treatment at steps 3 and 4 of NHLBI guidelines 78% receiving LABA 91% with at least 1 emergency room visit in previous year FEV ₁ % pred.: range, 15–139; median = 71 Mean Wasserfallen asthma symptom score = 17.3

		Study Population					
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)			
Bousquet et al. Predicting response to omalizumab, an anti-IgE antibody, in patients with allergic asthma. Chest 2004;125(4): 1378–1386. (Novartis Pharma AG and Genentech, Inc.) Djukanovic et al. Effects of treatment with anti-immunoglobulin E antibody omalizumab on airway inflammation in allergic asthma. Am J Respir Crit Care Med 2004; 170(6): 583–593.	Multicenter, double-blind, randomized, placebo-controlled Phase III study [combined data from Busse et al. (2001) and Solèr et al. (2001)] Multicenter, randomized, double- blind, placebo-controlled, parallel- group design (5 centers)	1,070 (1,070) 45 (42)	Age 12–76 yr; mean = 39.4 Gender 43% male, 57% female Ethnicity Not reported Total serum IgE, 30–860 IU/mL (mean = 197); 41% with history of emergency asthma treatment in past year Age 19–48 yr, median = 26 yr Gender 47% male, 53% female Ethnicity Not reported	Symptomatic allergic asthma with daily doses of BDP (200–2,000 mcg/day; mean = 725 mcg/day,) Duration,1–68 yr, mean = 20.6 yr FEV ₁ % pred., mean = 69; 21.5% with FEV ₁ \leq 65% pred. Stable asthma: 30 (66.7%) with mild asthma, 15 (33.3%) with moderately severe asthma Duration >1 yr Treatment with inhaled beta ₂ -agonists only No acute exacerbations for at least 6 weeks Positive skin-prick test for at least 1 of the following allergens: house-dust mite, cockroach, dog, or cat PC ₂₀ value, <8 mg/mL			
Holgate et al. Efficacy and safety of a recombinant anti-immunoglobulin E antibody (omalizumab) in severe allergic asthma. Clin	Multicenter, randomized, double- blind, placebo-controlled trial	246 (22 dropouts; ITT analysis)	Age 12–75 yr, mean = 40.8 yr	Sputum eosinophilia >2% or more of total nonsquamous cells Severe asthma requiring ≥1,000 mcg/day fluticasone for symptom control and positive SPTs to aeroallergen/s			
Exp Allergy 2004;34(4):632–638. (Novartis Pharma AG and Genentech Inc.)			Gender 39% male, 61% female Ethnicity Not reported Total serum IgE, 20–700 IU/mL	FEV ₁ % pred., mean = 64			

		Study Population					
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)			
Noga et al. Immunological and clinical changes in allergic asthmatics following treatment with omalizumab. Int Arch Allergy Immunol 2004;131(1): 46–52.	Substudy of a multicenter, randomized, double-blind, placebo- controlled, parallel-group trial	35 (35)	Age 23–61 yr, mean = 37.5 yr Gender 54% male, 46% female Ethnicity Not reported	Moderate-to-severe allergic asthma Duration ≥1 yr FEV ₁ % pred.: median = 79; range, 41–108 IgE mean = 165 IU/L All had positive skin-prick test to at least 1 of the tested allergens: house-dust mite, 76%; cat dander, 79%; dog dander, 57% ICS equivalent to BDP 500–1,000 mcg/day for at least 2 months Reversibility of >12% in FEV ₁ over baseline within 30 min after taking 200 mcg of salbuterol			
Vignola et al. Efficacy and tolerability of anti- immunoglobulin E therapy with omalizumab in patients with concomitant allergic asthma and persistent allergic rhinitis: SOLAR. Allergy 2004;59(7):709–717. (Novartis Pharma AG and Genentech Inc.)	Multicenter, randomized, double-blind, placebo-controlled, parallel-group trial	405 (ITT analysis; 20 withdrew)	Age 12–74 yr, mean = 38.4 yr Gender 45% male, 55% female Ethnicity Not reported	Moderate-to-severe allergic asthma (GINA) and persistent allergic rhinitis; 90% severe persistent; ≥400 mcg/day budesonide 39% receiving LABA FEV ₁ % pred., mean = 78.1			
Humbert et al. Benefits of omalizumab as add-on therapy in patients with severe persistent asthma who are inadequately controlled despite best available therapy (GINA 2002 step 4 treatment): INNOVATE. Allergy 2005;60(3):309–316.	Multicenter, randomized, double-blind, placebo-controlled study 18 centers in 14 countries	482 (419 for analysis; 52 withdrew)	Age 12–79 yr, mean = 43.3 yr Gender 34% male, 67% female Ethnicity 78% Caucasian, 7% Black, 15% other Serum IgE: range, 21–632 IU/mL; mean = 193 IU/mL	Severe persistent asthma FEV ₁ % pred.: mean = 61, range 18–101 PEF: mean = 305; range, 93–635 Rescue medications: mean = 6.1 puffs/day ICD 900–8,000 mcg/day, mean = 2,330 mcg/day All receiving ICS plus LABA 67% at high risk for asthma mortality			

		Study Population					
Citation (Sponsor)	Study Design	Study N (Number Evaluable)	Population Characteristics	Asthma Severity at Baseline (If Reported)			
Niebauer et al. Impact of omalizumab on quality-of-life outcomes in patients with moderate-to-severe allergic asthma. Ann Allergy Asthma Immunol 2006;96(2):316–326. (Genentech, Inc.)	Meta-analysis of randomized clinical trials that measured asthma-related quality of life using the Juniper Asthma Quality of Life Questionnaire (AQLQ)	5 trials, 2,056 subjects	Age 12–75 yr, mean = 39–42 yr (4 trials, n=1,722); 1 pediatric trial (n=334) mean = 9.4 yr Gender 70% male, 30% female in pediatric trial; 44% male, 56% female in 4 adolescent/adult trials Ethnicity Not reported	Moderate-to-severe allergic asthma (3 trials with adult and adolescent patients), allergic asthma (1 trial with n=334 children and adolescents), and asthma and allergic rhinitis (1 trial with adult and adolescent patients)			

n	Study Characteristics						
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Boulet et al. Inhibitory effects of an anti-IgE antibody E25 on allergen- induced early asthmatic response. Am J Respir Crit Care Med 1997;155(6): 1835– 1840. (Genentech Inc.)	Purpose/Objecti rhuMAb-E25 and response of inhale Arm 1 RhuMAb-E25 Arm 2 Placebo	ve: To document safet determine if it reduces ed aeroallergens 0.4 mL/kg on day 0; 0.2 mL/kg on days 7, 14, 28, 42, 56, and 70 (2.0 mg/kg IV)	y and tolerance of the early asthmatic 10 weeks treatment, 1 week off-treatment followup			Mean serum-free IgE decreased 89% after rhuMAb-E25 (p <0.001); no change occurred in placebo group. Allergen PC ₁₅ improved significantly after rhuMAb-E25 but not after placebo (p <0.002). Median change of 2.7 doubling doses	*Safety: 1 withdrawal after first dose, probably related to study drug *High BDP dose (≥800 mcg/day), history of emergency asthma treatment in past year, and FEV ₁ ≤65% pred. were predictive of greater probability of response.
Fahy et al. The effect of an anti-IgE monoclonal antibody on the early- and late- phase responses to allergen inhalation in asthmatic subjects. Am J Respir Crit Care Med 1997;155(6): 1828–1834. (Genentech Inc.)	Purpose/Objecti IgE on allergic air Arm 1 rhuMAb-E25 (n=10; 1 withdrawal) Arm 2 Placebo (n=9)	ve: To examine the eff way responses 5 mg/mL by 5-min IV (0.1 mL/kg) 150 mM NaCl, 10 mM acetate, pH 5.2 by 5-min IV	ects of neutralizing 9 weeks treatment, 1 week followup		*Trend was in favor of omalizumab for difference in FEV ₁ . RhuMAb-E25 significantly attenuated both early- and late- phase responses to airway challenge with allergen; reduced mean FEV ₁ from 30% to 18.8% vs. 33% to 34% (p=0.01) during early phase and from 24% to 9% vs. 20% to 18% during late response (p=0.047)	PC ₂₀ for methacholine improved, but not significantly. Free IgE concentrations in serum decreased significantly in rhuMAb-E25 group as compared to placebo group (p <0.001).	1 subject was withdrawn from rhuMAb-E25 treatment at 4 weeks because of asthma exacerbation.

		Study Characteristi	CS		Findings			
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other	
Milgrom et al. Treatment of allergic asthma with monoclonal anti-IgE antibody. N Engl J Med 1999;341(26): 1966–1973.	Purpose/Objecti as a treatment for Arm 1 High dose (HD) rhuMAb-E25 (n=106; 103 >4 weeks)	ve: To examine the ef allergic asthma 5.8 mcg/kg/ng IgE/mL	ficacy of rhuMAb-E25 20 weeks: 12 weeks of continued ICS and half dose of treatment on days 0 and 4, full dose on	After 12 weeks, albuterol use reduced by 1.8 puffs/day in HD group (p=0.02) and by 1.2 puffs/day in LD group (p=0.24) vs. 0.8 puffs/day in placebo group. Decreases were maintained at 20 weeks.	PEF favored treatment at all points ($p < 0.01$); FEV ₁ % pred. was significantly higher for treatment group between weeks 4 and 12	30% HD (p=0.03) and 28% LD groups (p=0.01) had exacerbations vs. 45% of placebo group. *Asthma symptom score at 12 weeks: 2.8 for HD group and 2.8 for LD group vs. 3.1 for placebo group (p=0.008 and p=0.005, respectively)	3 HD, 3 LD, and 8 placebo subjects withdrew due to AE.	
(Genentech Inc.)	≥4 weeks) Arm 2 Low dose (LD) rhuMAb-E25 (n=106; 103 ≥4 weeks) Arm 3 Placebo (n=105; 100 ≥4 weeks)	2.5 mcg/kg/ng IgE/mL	day 7 and then once every 2 weeks; For 8 weeks, treatment continued and ICS was tapered. 10 weeks of followup		weeks 4 and 12 and between weeks18 and 28. Morning PEF increased 30.7 L/min in HD group (p=0.007) and 18.6 L/min in LD group vs. 11.3 L/min in placebo group. At 20 weeks, increase from baseline was 29.9 L/min in HD group (p=0.02), 20.8 L/min LD group (p=0.046), and 10.2 L/min in placebo group.			

		Study Characterist	ics		Findings		
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Busse et al. Omalizumab, anti- IgE recombinant humanized monoclonal antibody, for the treatment of severe allergic Asthma. J Allergy Clin Immunol 2001;108(2): 184–190. (Novartis Pharmaceuticals Corp. and Genetech, Inc.)	Purpose/Objecti subcutaneous on severe allergic as controlled with IC Arm 1 Omalizumab (n=268; 19 withdrawals) Arm 2 Placebo (n=257; 34 withdrawals)	ve: To assess the efficient nalizumab in adolescent sthma whose disease w S 0.016 mg/kg IgE (IU/mL) every 4 weeks for 16 weeks; approximately 25% reduction every 2 weeks for weeks 16–28 Rescue albuterol, maximum of 8 puffs/day was allowed.	cacy and tolerability of ts and adults with vas not adequately 28 weeks (16-week stable phase, 12-week reduction phase)	Median reduction in ICS dose was greater in omalizumab group (75% vs. 50%, p <0.001). ≥50% reduction in BDP dose for 72.4% of omalizumab group vs. 54.9% of placebo group (p <0.001) BDP was discontinued in 39.6% of omalizumab and 19.1% of placebo group (p <0.001).		*Stable phase: 14.6% of intervention vs. 23.3% of placebo group experienced exacerbations (p=0.009), with mean of 0.28 vs. 0.54 (p=0.006) exacerbations, respectively. Reduction phase: 21.3% of intervention vs. 32.3% of placebo group experienced exacerbations (p=0.004), with average of 0.39 vs. 0.66 (p=0.003) exacerbations. Total IgE increased in treatment group and did not change in placebo group.	At 28 weeks, 60.6% of treatment vs. 28.1% of placebo patients rated treatment effectiveness as good or excellent (p <0.001).

		Study Characteristi	cs		Findings			
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other	
Milgrom et al. Treatment of childhood asthma with anti- immunoglobulin E antibody (omalizumab). Pediatrics 2001;108(2):E36. (Genentech, Inc.; Novartis Pharmaceuticals Corp.)	Purpose/Objecti sparing effect of o exacerbations in asthma who requind Arm 1 Omalizumab (O) (n=225; n=209 completers) Arm 2 Placebo (P) (n=109; n=97 completers)	ve: To evaluate the sa omalizumab and its imp children with moderate ire daily treatment with Subcutaneous dose, based on body weight and initial minimum of 0.016 mg/kg/IgE (IU/mL), per 4 weeks, was based on dosing chart for 16 weeks; dose was tapered approximately 25% of baseline dose every 2 weeks until elimination or worsening of asthma.	fety and steroid- bact on asthma to severe allergic ICSs 16-week stable- steroid phase and 12 weeks of steroid dose-reduction phase after 4- to 6-week run-in phase All children switched from ICS to equivalent dose of BDP for asthma control. Salbutamol 2 puffs (90 mcg/puff ex mouthpiece equal to 100 mcg/puff ex valve) was used as needed, with maximum 8 puffs/day, for rescue medication.	Greater proportion of O reduced ICS dose vs. P (p=0.001). Median reduction in dose was 100% in O vs. 66.7% in P. BDP withdrawal was completed in 55% of O group vs. 39% of P group (p=0.004).	Little change occurred in morning PEF, FEV ₁ , FVC, or FEF _{25%-75%} during either phase, with minimal difference between groups.	During treatment phase, incidence of exacerbations was lower for O vs. P group (18.2% vs. 38.5%, p <0.001), and number of episodes/patient was lower for O vs. P group (0.42 vs. 2.72, p <0.001). Mean duration of episodes was similar in groups during both phases (range, 10–14 days). Fewer subjects in O vs. P group required urgent unscheduled physician visit (12.9% vs. 30.3%, p=0.001). At week 28, median number of puffs/day of rescue medication was 0 in O group and 0.46 in P group; change from baseline favored O treatment (p=0.004).	No treatment-related SAE occurred. Drug-related AEs were more frequent in O vs. P subjects (6.2% vs. 0.9%, p=0.029). Investigators' global evaluation of effectiveness favored O vs. P treatment (p <0.001): excellent for 31.5% of O group vs. 16.3% of P group and good for 44.7% of O group vs. 32.7% of P group. Fewer missed school days occurred for O vs. P subjects (0.65 vs. 1.21 days, p=0.040).	

		Study Characteristics			Findings		
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Soler et al. The anti-IgE antibody omalizumab	Purpose/Objecti corticosteroid-spa subcutaneously ir	ve: To evaluate the ef aring effect of omalizum a allergic asthma	ficacy, safety, and nab administered	Proportion able to reduce BDP dose was higher at end of steroid-reduction phase compared to stable-steroid	FEV ₁ was significantly higher for omalizumab	*Omalizumab group had 58% fewer exacerbations per patient vs. placebo group during stable-steroid phase	Suspected drug-related AE events for 1.1% of omalizumab group; none were serious.
reduces exacerbations and steroid requirement in allergic asthmatics. Eur Respir J 2001;18(2): 254–261. Novartis Pharma AG and Genentech Inc.)	Arm 1 Omalizumab (n=274; 19 withdrawals) Arm 2 Placebo (n=232; 40 withdrawals)	0.016 mg/kg IU IgE/mL every 4 weeks Rescue salbutamol of 100 mcg/puff was allowed throughout the study.	28 weeks, after 4- to 6-week run-in 16-week stable steroid phase, then 8-week reduction phase, and lowest dose held for 4 weeks	phase higher in omalizumab group (p <0.001). Reduction in BDP <u>></u> 50% was achieved for 79% of omalizumab group and 55% of placebo group.	than placebo patients between weeks 18 and 28 (ITT analysis). PEF values favored omalizumab at all time points of the stable-steroid phase (ITT analysis).	(p <0.001) and 52% fewer exacerbations during steroid-reduction phase (p <0.001).	

		Study Characteristi	ics		Findings		
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Walker et al. Anti- IgE for chronic asthma. Cochrane Database Syst Rev	Purpose/Objecti studies that have placebo or other of chronic asthma	ve: To compare the cli compared anti-IgE mo conventional therapy in	inical outcomes in noclonal antibodies to the treatment of	Subcutaneous O vs. P during steroid- reduction phase: Achieving complete ICS withdrawal, OR 2.50, 95% CI 2.00 to 3.13 (4 trials, n=1,534); achieving	Subcutaneous O vs. P during stable steroid phase: No difference found in	Subcutaneous O vs. P during stable- steroid phase: Number with exacerbations, OR 0.46, 95% CI 0.35 to 0.61 (3 trials, n=1,405); asthma	Reductions in serum free IgE ranged from 89% to 99% in all trials, despite different dosing regimens.
2002;(3): CD003559. (Garfield Weston Foundation UK ; NHS Research and Development UK ; The Thriplow Charitable Trust UK)	Arm 1 Omalizumab (O) Arm 2 Placebo (P)	Intravenous route: 5.8 mcg/kg/ng IgE/mL, 2.5 mcg/kg/ng IgE/mL Inhaled route: 1 mg or 10 mg Subcutaneous route: 0.016 mg/kg/IU/mL	12–16 weeks (stable-steroid phase) followed by steroid-reduction phase Extension phase (2 trials)	1 >50% reduction in ICS usage, OR 2.50, 95% CI 2.02 to 3.10 (4 trials, n=1645)	FEV ₁ (2 trials, n=1,071) or PEF (4 trials, n=1,651). <i>Intravenous O vs.</i> <i>P</i> : No difference found in FEV ₁ or PEF (2 trials, n=39).	exacerbations/patient, diff. -0.19 , 95% CI -0.29 to -0.09 (4 trials, n=1,651); rescue puffs/day, diff. -0.73 , 95% CI -10.7 to -0.39 (2 trials, n=1,071); symptom scores diff. -0.48 , 95% CI -0.67 to -0.28 (2 trials, n=1,071). Subcutaneous O vs. P during steroid- reduction phase: Number with exacerbation, OR 0.46, 95% CI 0.36 to 0.59 (3 trials, n=1,388); exacerbations requiring hospitalization, OR 0.11, 95% CI 0.03 to 0.48 (3 trials, n=1,405); exacerbations/patient, diff. -0.27 , 95% CI -0.38 to -0.17 (4 trials, n=1,634); rescue puffs/day, diff. -0.73 , 95% CI -1.06 to -0.40 (2 trials, n=1,071) Intravenous O vs. P: No difference in rescue medication use 1 week after end of treatment or in symptom scores (2 trials, n=39)	

	Study Characteristics				Findings		
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Corren et al. Omalizumab, a recombinant humanized anti-IgE	Purpose/Objecti (Xolair) a recomb antibody, on the r term therapy	ve: To investigate the inant humanized mono ate of serious exacerba	effect of omalizumab clonal anti-IgE ations during long-			*Incidence rate of unscheduled visits was lower for O group vs. P group (21.3 vs. 35.5 per 100 patient-yr; RR 0.60, 95% CI 0.44 to 0.81, p <0.01).	
antibody, reduces asthma-related emergency room visits and hospitalizations in patients with allergic asthma. J Allergy Clin Immunol 2003; 111(1):87–90. (Novartis Pharma AG; Genentech, Inc.)	Arm 1 Omalizumab (O) (n=767) Arm 2 Placebo (P) (n=638)	Subcutaneous injection every 2 or 4 weeks, dosed according to body weight and baseline IgE (≥0.016 mg/kg/IgE [IU/mL] every 4 weeks)	12 months. Dosages of BDP were stable over initial 16 weeks (steroid-stable phase) and reduced over 8 weeks (25% every 2 weeks), with lowest effective dose maintained for 4 weeks (steroid- reduction phase). Minimum effective dose was maintained for 24 weeks (extension phase).			 *Incidence rate of ED treatment for exacerbation was lower for O vs. P group (1.8 vs. 3.8 per 100 patient-yr; RR 0.47, 95% Cl 0.24 to 1.01, p=0.05). *Incidence rate of asthma-related hospitalization was 92% lower in O vs. P group (0.26 vs. 3.42 per 100 patient-yr, RR 0.08, 95% Cl 0 to 0.25, p <0.01). Mean number of days per asthma- related hospitalization for O group was less than for P group (2.00 vs. 5.39, p=0.15). 	

		Study Characteristi	CS		Findings		
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Berger et al. Evaluation of long- term safety of the anti-IgE antibody, omalizumab, in children with allergic asthma.	Purpose/Objective : To evaluate the long-term effects of the anti-lgE antibody omalizumab in children with asthma treated for full 52 weeks				FEV ₁ remained stable	81.4% did not require other medication. No change occurred in mean BDP	*Adverse events: 93% experienced AE unrelated to drug; 6.7% AE were suspected as related to drug
	Arm 1 Omalizumab (n=225)	0.016 mg/kg/lgE (IU/mL) per 4 weeks	28-week treatment, 24-week open-label extension, 12 weeks off study drug.			dosage No anti-omalizumab antibodies were detected; no clinically significant changes occurred in vital signs.	(1 SAE resolved in 20 minutes).
Immunol 2003;91(2): 182–188. (Novartis Pharmaceuticals Corp.)	Arm 2 Placebo (not included in this report)					Log ₁₀ methacholine PC ₂₀ was significant (p <0.05) after treatment (change of 0.9 doubling doses); no change was found after placebo.	
Ayres et al. Efficacy and tolerability of anti-immunoglobulin E therapy with omalizumab in patients with poorly	Purpose/Objective : To investigate the efficacy and tolerability of omalizumab in patients with poorly controlled allergic asthma in an open-label study in which omalizumab was added to and compared against best standard care (BSC) as defined by the NHLBI		Patients treated with omalizumab $(n=173)$ reduced mean daily dose of ICS (-342 mcg/day), and those with BSC alone showed slight increase $(+68 mcg/day)$, p <0.001.	Significant difference in FEV ₁ of 2.48 L for omalizumab vs. 2.28 L for BSC	*Those treated with omalizumab experienced 4.84 fewer asthma deterioration-related incidents (ADRI) vs. those treated with BSC alone (49.6% reduction); 36.1% vs. 20.2%,	Percentage of AE was not significantly different between groups (p=0.116). 48 patients (16.5% of omalizumab group and 13.2% of BSC group) experienced SAE during study.	
controlled (moderate-to- severe) allergic asthma. Allergy 2004;59(7): 701–708.	Arm 1 BSC with omalizumab (n=206) Arm 2 BSC without omalizumab (n=106)	0.016 mg/kg/IgE (IU/mL) per 4 weeks Rescue salbutamol was permitted throughout the study.	52 weeks		aione, p=0.02	during the study. Median time to first ADRI was 126 vs. 75 days, respectively (p=0.03). Lower annualized mean number of exacerbations with occurred with omalizumab than BSC alone (1.12 vs. 2.86, p <0.001). 49.5% omalizumab subjects vs. 26.4% BSC only were exacerbation free (p=0.001).	

	Study Characteristics				Findings		
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Bousquet et al. Predicting response to omalizumab, an anti-IgE antibody, in patients with allergic asthma. Chest 2004;125(4): 1378–1386. (Novartis Pharma AG and Genentech, Inc.)	Purpose/Objecti predictive of best asthma, time to o need to be treated predicted Arm 1 Omalizumab (n=542) Arm 2 Placebo (n=528)	ve: To determine base response to omalizuma nset of response, and h d before a response co 0.016 mg/kg/IgE subcutaneously every 4 weeks in addition to stable BDP therapy Rescue BDP and albuterol were permitted.	line characteristics ab therapy for allergic now long patients uld be accurately 16 weeks		Significant improvements in FEV ₁ in the treatment group as compared to placebo group were maintained for the entire study (p values ranged from <0.001 to 0.019).	Time to exacerbation was longer for omalizumab patients (p <0.001). Probability of exacerbation by week 16 was 30% for placebo subjects and 16% for omalizumab subjects.	Odds of being responder were 2.25 times higher (95% CI, 1.68 to 3.01) with omalizumab than with placebo. 61% of responders at 16 weeks had responded at 4 weeks and 87% had responded by 12 weeks.

		Study Characteristi	cs		Findings		
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Djukanovic et al. Effects of treatment with anti-	Purpose/Objective : To determine whether omalizumab has anti-inflammatory effects in the airways of patients with allergic asthma				No difference in change in airway responsiveness	*Mean percent of eosinophils in induced sputum decreased from 6.6% to 1.7% for O group (p <0.001) and	
immunoglobulin E antibody omalizumab on airway inflammation in allergic asthma. Am J Respir Crit Care Med 2004;170(6): 583–593.	Arm 1 Omalizumab (O) (n=22; n=21 completers) Arm 2 Placebo (P) (n=23; n=22 completers)	150–300 mg every 4 weeks or 225–375 mg every 2 weeks on the basis of concentration of serum total IgE and body weight at baseline	16 weeks after run-in period of 3 weeks		(p=0.14) was found between groups. In O group, PC ₂₀ changed from 1.01 to 0.73 mg/mL (p=0.47); in P group PC ₂₀ changed from 0.54 to 0.67 (p=0.26).	from 8.5% to 7.0% for P group (p >0.05). Difference in change between groups was -4.6% (p=0.05). Between-group differences for O vs. P for were: eosinophils in submucosa (p=0.03); IgE ⁺ cells in epithelium (p=.001) and submucosa (p <0.001); CD3 ⁺ (p=0.01), CD4 ⁺ (p=0.005), and CD8 ⁺ (p=0.05); T lymphocytes and B lymphocytes (p=0.02).	
Holgate et al. Efficacy and safety of a recombinant anti-immunoglobulin E antibody	Purpose/Objective:To evaluate the ability of omalizumab to improve disease control to enable ICS reduction in patients with severe allergic asthmaArm 10.016 mg/kg/lgE (IU/mL) every32 weeks, after 6–10 week run-in period		*Patients receiving omalizumab had greater mean reduction in fluticasone dose as compared to those receiving placebo (57.2% vs. 43.3%, p=0.003), with 73.8% vs. 50.8% achieving \geq 50% dose reduction (p=0.001).	Trend in favor of omalizumab for FEV ₁ throughout; trend was significant at weeks 4, 20, 28	Difference in reduction of exacerbation rates did not reach statistical significance. 58% of omalizumab vs. 39% of placebo patients had improvement in asthma rolated Oct. (p. c0.01)	n 1 omalizumab and 5 placebo patients had SAE not considered drug related. Completer rates were 91.3% for omalizumab and 90.8% for placebo group.	
severe allergic asthma. Clin Exp Allergy 2004;34(4): 632–638. (Novartis Pharma AG and Genentech	(n=126; 115 completed) Arm 2 Placebo (n=120;	4 weeks Beta ₂ -agonists were allowed as needed.	followed by 16-week corticosteroid- reduction phase	Median reduction in prednisone was 50% in high dose (p=0.045) and 65% in low dose (p=0.11) groups vs. 0% in placebo. 21.4% vs. 15.0% with 100% reduction (p=0.198)	and 30.		

	Study Characteristics				Findings		
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Noga et al. Immunological and clinical changes in allergic asthmatics following treatment with omalizumab. Int Arch Allergy Immunol 2004; 131(1): 46–52.	Purpose/Objecti using more sensit define factors and cellular reactions mode of action of Arm 1 Omalizumab (O) (n=18) Arm 2 Placebo (P) (n=17)	ve: To strengthen prev ive parameters of lung a mediators involved in in order to improve our omalizumab Subcutaneous administration of at least 0.016 mg/kg/IgE every 4 weeks	vious findings by function and to immunological and understanding of the 16 weeks after 6-week run-in Stable dose of BMD 500–1,200 mcg/day		Eosinophils decreased in O vs. P subjects (6.1% to 1.3% vs. 3.5% to 6.0%; p < 0.01). R _{aw} decreased in O v. P group ($p < 0.01$) but returned to baseline 3 months after discontinuation of treatment. PC ₂₀ increased in O vs. P subjects ($p < 0.01$), but after discontinuation of therapy there was no difference. No changes occurred in FEV ₁ in either group. Free IgE levels decreased ($p < 0.01$) to <10 IU/mL in all O subjects vs. no change in P subjects.		Interleukin-13 (IL-13) levels decreased in O vs. P subjects (9.4 to 7.0 pg/mL vs. 7.2 to 8.5 pg/mL; p <0.01). No difference was found for IL-6, IL-10, and s-ICAM levels. Decrease in IL-5 and IL-8 in the O group did not reach significance. Area of wheal reaction decreased in O group for all 3 allergens (p <0.001).

		Study Characteristi	CS		Findings		
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Vignola et al. Efficacy and tolerability of anti- immunoglobulin E therapy with omalizumab in patients with concomitant allergic rhinitis: SOLAR. Allergy 2004;59(7): T09–717.	Purpose/Objective: To evaluate the efficacy and safety of opmalizumab in patients with concomitant asthma and persistent allergic rhinitis			Treatment group difference occurred in FEV ₁	*20.6% of omalizumab vs. 30.1% of placebo (p=0.02) experienced exacerbations (using imputed values),	*Greater proportion treated with omalizumab had ≥1 point improvement in asthma QoL (57.5% vs. 40.6%, p <0.001).	
	Arm1 Omalizumab (n=209; 5 withdrawals) Arm 2 Placebo (n=196; 15 withdrawals)	0.016 mg/kg IU IgE/mL every 4 weeks	28 weeks after 4-week run-in		(73 mL, p=0.032), FVC (84 L, p=0.016), and PEF (11 L/min, p <0.001) compared with placebo.	and 18.2% vs. 25.5% experienced exacerbations (p=0.055) without imputed values. Rate of exacerbations was 0.25 vs. 0.40 (p=0.02).	SAE rates were similar (1.4% vs. 1.5%) in the 2 groups.
(Novartis Pharma AG and Genentech Inc.)							

		Study Characteristi	cs		Findings			
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other	
Humbert et al. Benefits of omalizumab as add-on therapy in patients with severe persistent asthma who are inadequately controlled despite best available therapy (GINA 2002 step 4 treatment): INNOVATE. Allergy 2005;60(3): 309– 316.	Purpose/Objecti omalizumab on a persistent asthma GINA step 4 thera Arm 1 Omalizumab (n=209) Arm 2 Placebo (n=210)	ve: To evaluate the ef sthma exacerbations in a who were inadequate apy 0.016 mg/kg/lgE (IU/mL) every 4 weeks	fect of add-on patients with severe y controlled despite 28 weeks after 8-week run-in phase (16-week followup phase not reported here)	Difference in rescue medication use was not significant.	Change from baseline in mean morning PEF was greater for omalizumab than for placebo ($p=0.042$). Improvement in FEV ₁ was 190 mL for omalizumab and 96 mL for placebo ($p=0.043$).	*Exacerbation rate for omalizumab group was 0.68 vs. 0.91 for placebo group (p=0.042, rate ratio 0.738) after adjusting for baseline differences in history (NNT=2.2) Severe exacerbation rate was lower in omalizumab vs. placebo (0.24 vs. 0.48; p=0.002) (NNT=2.2). Mean change in symptom score was greater with omalizumab vs. placebo treatment (p=0.039).	Asthma QoL improvement from baseline of ≥0.5 points occurred for 60.8% of omalizumab group vs. 47.8% of placebo group (p=0.008).	

		Study Characteristi	CS		Findings		
Citation (Sponsor)	Treatment	Dose	Duration of Active Treatment; Duration of Postintervention/ Off-Treatment Followup	Taper/ Decrease Steroids	Lung Function	Exacerbations/ Symptoms	Other
Niebauer et al. Impact of omalizumab on quality-of-life outcomes in patients with moderate-to-severe	Purpose/Objectiv outcomes associa to-severe allergic Arm 1 Omalizumab (O)	e: To summarize asthr ated with omalizumab t asthma Not reported Patients were permitted to use	ha-related QoL herapy in moderate- 4- to 6-week run-in period, 16-week steroid-stabilization	All results refer to Juniper Asthma Quality of Life Questionnaire (AQLQ) Overall effect sizes during steroid- reduction phase were 1.73 for O and 1.31 for P groups. <i>Change >0.5 in AQLQ for O vs. P:</i> steroid-stabilization phase_OR 1.35			
allergic asthma. Ann Allergy Asthma Immunol 2006;96(2): 316–326. (Genentech, Inc.)	Arm 2	albuterol metered- dose inhaler as needed (5 trials), treated concomitantly with BDP (3 trials), used fluticasone (1 trial), or used budesonide Turbohaler (1 trial).	pnase, 12- to 16-week steroid- reduction phase, and either an open-label or double-blind extension phase. 2 trials lasted 52 weeks, 1 lasted 32 weeks, and 2 lasted 28 weeks.	95% CI 1.11 to 1.64 (4 trials, n=1,649); steroid-reduction phase, OR 1.69, 95% CI 1.40 to 2.05 (5 trials, n=1,864); extension phase, OR 1.50, 95% CI 1.15 to 1.95 (3 trials, n=1,078) <i>Change</i> >1.0 in AQLQ for O vs. P: steroid-stabilization phase, OR=1.61, 95% CI 1.29 to 2.00 (4 trials, n=1,649); steroid-reduction phase, OR 2.03, 95% CI 1.66 to 2.47 (5 trials, n=1,864); extension phase, OR 1.25.			
	Placebo (P)			95% Cl 0.97 to 1.59 (3 trials, n=1,078) with evidence of heterogeneity (p=0.01). <i>Change</i> >1.5 in AQLQ for O vs. P: steroid-stabilization phase, OR1.80, 95% Cl 1.36 to 2.38 (4 trials, n=1,649); steroid-reduction phase, OR 2.11, 95% Cl 1.68 to 2.65 (5 trials, n=1864); extension phase, OR 1.59, 95% Cl 1.21 to 2.08 (3 trials, n=1,078) with evidence of heterogeneity (p=0.01)			