

**HIGHLIGHTS OF PRESCRIBING INFORMATION**

These highlights do not include all the information needed to use BENLYSTA safely and effectively. See full prescribing information for BENLYSTA.

**BENLYSTA® (belimumab)**  
**for injection, for intravenous use only**  
**Initial U.S. Approval: 2011**

**RECENT MAJOR CHANGES**

Warnings and Precautions (5.4) 3/2012

**INDICATIONS AND USAGE**

BENLYSTA is a B-lymphocyte stimulator (BLyS)-specific inhibitor indicated for the treatment of adult patients with active, autoantibody-positive, systemic lupus erythematosus who are receiving standard therapy. (1, 14)

**Limitations of Use:** The efficacy of BENLYSTA has not been evaluated in patients with severe active lupus nephritis or severe active central nervous system lupus (1). BENLYSTA has not been studied in combination with other biologics or intravenous cyclophosphamide (1). Use of BENLYSTA is not recommended in these situations.

**DOSAGE AND ADMINISTRATION**

- Recommended dosage regimen is 10 mg/kg at 2-week intervals for the first 3 doses and at 4-week intervals thereafter. Reconstitute, dilute and administer as an intravenous infusion only, over a period of 1 hour. (2.1)
- Consider administering premedication for prophylaxis against infusion reactions and hypersensitivity reactions (2.2)

**DOSAGE FORMS AND STRENGTHS**

Single-use vials of belimumab lyophilized powder:

- 120 mg per vial (3)
- 400 mg per vial (3)

**CONTRAINDICATIONS**

Previous anaphylaxis to belimumab. (4)

**WARNINGS AND PRECAUTIONS**

- Mortality:** There were more deaths reported with BENLYSTA than with placebo during the controlled period of clinical trials. (5.1)
- Serious Infections:** Serious and sometimes fatal infections have been reported in patients receiving immunosuppressive agents, including BENLYSTA. Use with caution in patients with chronic infections. Consider interrupting BENLYSTA therapy if patients develop a new infection during BENLYSTA treatment. (5.2)
- Hypersensitivity Reactions, Including Anaphylaxis:** Serious and fatal reactions have been reported. BENLYSTA should be administered by healthcare providers prepared to manage anaphylaxis. Monitor patients during and for an appropriate period of time after administration of BENLYSTA. (2.2, 5.4)
- Depression and suicidality** have been reported in BENLYSTA studies. Patients should be instructed to contact their healthcare provider if they experience new or worsening depression, suicidal thoughts or other mood changes. (5.6)
- Immunization:** Live vaccines should not be given concurrently with BENLYSTA. (5.7)

**ADVERSE REACTIONS**

Common adverse reactions (≥5%) in clinical trials were: nausea, diarrhea, pyrexia, nasopharyngitis, bronchitis, insomnia, pain in extremity, depression, migraine, and pharyngitis. (6.1)

**To report SUSPECTED ADVERSE REACTIONS, contact Human Genome Sciences, Inc. at 1-877-423-6597 or FDA at 1-800-FDA-1088 or [www.fda.gov/medwatch](http://www.fda.gov/medwatch).**

**USE IN SPECIFIC POPULATIONS**

- Pregnancy:** Registry available. (8.1)

See 17 for **PATIENT COUNSELING INFORMATION** and Medication Guide.

Revised: March 2012

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\* Sections or subsections omitted from the full prescribing information are not listed

1 **FULL PRESCRIBING INFORMATION**

2 **1 INDICATIONS AND USAGE**

3 BENLYSTA<sup>®</sup> (belimumab) is indicated for the treatment of adult patients with active,  
4 autoantibody-positive, systemic lupus erythematosus (SLE) who are receiving standard therapy.

5

6 *Limitations of Use*

7 The efficacy of BENLYSTA has not been evaluated in patients with severe active lupus nephritis  
8 or severe active central nervous system lupus. BENLYSTA has not been studied in combination  
9 with other biologics or intravenous cyclophosphamide. Use of BENLYSTA is not recommended  
10 in these situations.

11 **2 DOSAGE AND ADMINISTRATION**

12

13 **2.1 Dosage Schedule**

14 BENLYSTA is for intravenous infusion **only** and must be reconstituted and diluted prior to  
15 administration [*see Dosage and Administration (2.3)*]. Do not administer as an intravenous push  
16 or bolus.

17

18 The recommended dosage regimen is 10 mg/kg at 2-week intervals for the first 3 doses and at  
19 4-week intervals thereafter. Reconstitute, dilute and administer as an intravenous infusion only,  
20 over a period of 1 hour. The infusion rate may be slowed or interrupted if the patient develops an  
21 infusion reaction. The infusion must be discontinued immediately if the patient experiences a  
22 serious hypersensitivity reaction [*see Contraindications (4), Warnings and Precautions (5.4)*].

23

24 **2.2 Premedication Recommendations**

25 Prior to dosing with BENLYSTA, consider administering premedication for prophylaxis against  
26 infusion reactions and hypersensitivity reactions. [*see Warnings and Precautions (5.4,5.5) and*  
27 *Adverse Reactions (6.1)*].

28

29 **2.3 Preparation of Solutions**

30 BENLYSTA is provided as a lyophilized powder in a single-use vial for intravenous infusion  
31 only and should be reconstituted and diluted by a healthcare professional using aseptic technique  
32 as follows:

33 **Reconstitution Instructions**

- 34 1. Remove BENLYSTA from the refrigerator and allow to stand 10 to 15 minutes for the vial to  
35 reach room temperature.
- 36 2. Reconstitute the BENLYSTA powder with Sterile Water for Injection, USP, as follows. The  
37 reconstituted solution will contain a concentration of 80 mg/mL belimumab.
- 38 • Reconstitute the 120 mg vial with 1.5 mL Sterile Water for Injection, USP.
  - 39 • Reconstitute the 400 mg vial with 4.8 mL Sterile Water for Injection, USP.
- 40 3. The stream of sterile water should be directed toward the side of the vial to minimize  
41 foaming. Gently swirl the vial for 60 seconds. Allow the vial to sit at room temperature  
42 during reconstitution, gently swirling the vial for 60 seconds every 5 minutes until the  
43 powder is dissolved. *Do not shake*. Reconstitution is typically complete within 10 to  
44 15 minutes after the sterile water has been added, but it may take up to 30 minutes. Protect

- 45 the reconstituted solution from sunlight.
- 46 4. If a mechanical reconstitution device (swirler) is used to reconstitute BENLYSTA, it should  
47 not exceed 500 rpm and the vial swirled for no longer than 30 minutes.
- 48 5. Once reconstitution is complete, the solution should be opalescent and colorless to pale  
49 yellow, and without particles. Small air bubbles, however, are expected and acceptable.

#### 50 **Dilution Instructions**

- 51 6. Dextrose intravenous solutions are incompatible with BENLYSTA. BENLYSTA should only  
52 be diluted in 0.9% Sodium Chloride Injection, USP. Dilute the reconstituted product to  
53 250 mL in 0.9% Sodium Chloride Injection, USP (normal saline) for intravenous infusion.  
54 From a 250-mL infusion bag or bottle of normal saline, withdraw and discard a volume equal  
55 to the volume of the reconstituted solution of BENLYSTA required for the patient's dose.  
56 Then add the required volume of the reconstituted solution of BENLYSTA into the infusion  
57 bag or bottle. Gently invert the bag or bottle to mix the solution. Any unused solution in the  
58 vials must be discarded.
- 59 7. Parenteral drug products should be inspected visually for particulate matter and discoloration  
60 prior to administration, whenever solution and container permit. Discard the solution if any  
61 particulate matter or discoloration is observed.
- 62 8. The reconstituted solution of BENLYSTA, if not used immediately, should be stored  
63 protected from direct sunlight and refrigerated at 2° to 8°C (36° to 46°F). Solutions of  
64 BENLYSTA diluted in normal saline may be stored at 2° to 8°C (36° to 46°F) or room  
65 temperature. The total time from reconstitution of BENLYSTA to completion of infusion  
66 should not exceed 8 hours.
- 67 9. No incompatibilities between BENLYSTA and polyvinylchloride or polyolefin bags have  
68 been observed.

#### 70 **2.4 Administration Instructions**

- 71 1. The diluted solution of BENLYSTA should be administered by intravenous infusion only,  
72 over a period of 1 hour.
- 73 2. BENLYSTA should be administered by healthcare providers prepared to manage  
74 anaphylaxis. [see *Warnings and Precautions (5.4)*]
- 75 3. BENLYSTA should not be infused concomitantly in the same intravenous line with other  
76 agents. No physical or biochemical compatibility studies have been conducted to evaluate the  
77 coadministration of BENLYSTA with other agents.

### 78 **3 DOSAGE FORMS AND STRENGTHS**

79 Single-use vials of belimumab lyophilized powder for injection:

- 80 • 120 mg per vial  
81 • 400 mg per vial

### 82 **4 CONTRAINDICATIONS**

83 BENLYSTA is contraindicated in patients who have had anaphylaxis with belimumab.

### 84 **5 WARNINGS AND PRECAUTIONS**

#### 86 **5.1 Mortality**

87 There were more deaths reported with BENLYSTA than with placebo during the controlled  
88 period of the clinical trials. Out of 2133 patients in 3 clinical trials, a total of 14 deaths occurred

89 during the placebo-controlled, double-blind treatment periods: 3/675 (0.4%), 5/673 (0.7%),  
90 0/111 (0%), and 6/674 (0.9%) deaths in the placebo, BENLYSTA 1 mg/kg, BENLYSTA 4  
91 mg/kg, and BENLYSTA 10 mg/kg groups, respectively. No single cause of death predominated.  
92 Etiologies included infection, cardiovascular disease and suicide.

93

## 94 **5.2 Serious Infections**

95 Serious and sometimes fatal infections have been reported in patients receiving  
96 immunosuppressive agents, including BENLYSTA. Physicians should exercise caution when  
97 considering the use of BENLYSTA in patients with chronic infections. Patients receiving any  
98 therapy for chronic infection should not begin therapy with BENLYSTA. Consider interrupting  
99 BENLYSTA therapy in patients who develop a new infection while undergoing treatment with  
100 BENLYSTA and monitor these patients closely.

101

102 In the controlled clinical trials, the overall incidence of infections was 71% in patients treated  
103 with BENLYSTA compared with 67% in patients who received placebo. The most frequent  
104 infections (>5% of patients receiving BENLYSTA) were upper respiratory tract infection,  
105 urinary tract infection, nasopharyngitis, sinusitis, bronchitis, and influenza. Serious infections  
106 occurred in 6.0% of patients treated with BENLYSTA and in 5.2% of patients who received  
107 placebo. The most frequent serious infections included pneumonia, urinary tract infection,  
108 cellulitis, and bronchitis. Infections leading to discontinuation of treatment occurred in 0.7% of  
109 patients receiving BENLYSTA and 1.0% of patients receiving placebo. Infections resulting in  
110 death occurred in 0.3% (4/1458) of patients treated with BENLYSTA and in 0.1% (1/675) of  
111 patients receiving placebo.

112

## 113 **5.3 Malignancy**

114 The impact of treatment with BENLYSTA on the development of malignancies is not known. In  
115 the controlled clinical trials, malignancies (including non-melanoma skin cancers) were reported  
116 in 0.4% of patients receiving BENLYSTA and 0.4% of patients receiving placebo. In the  
117 controlled clinical trials, malignancies, excluding non-melanoma skin cancers, were observed in  
118 0.2% (3/1458) and 0.3% (2/675) of patients receiving BENLYSTA and placebo, respectively. As  
119 with other immunomodulating agents, the mechanism of action of BENLYSTA could increase  
120 the risk for the development of malignancies.

121

## 122 **5.4 Hypersensitivity Reactions, Including Anaphylaxis**

123 Hypersensitivity reactions, including anaphylaxis and death, have been reported in association  
124 with BENLYSTA. Delay in the onset of acute hypersensitivity reactions has been observed.  
125 Limited data suggest that patients with a history of multiple drug allergies or significant  
126 hypersensitivity may be at increased risk. In the controlled clinical trials, hypersensitivity  
127 reactions (occurring on the same day of infusion) were reported in 13% (191/1458) of patients  
128 receiving BENLYSTA and 11% (76/675) of patients receiving placebo. Anaphylaxis was  
129 observed in 0.6% (9/1458) of patients receiving BENLYSTA and 0.4% (3/675) of patients  
130 receiving placebo. Manifestations included hypotension, angioedema, urticaria or other rash,  
131 pruritus, and dyspnea. Due to overlap in signs and symptoms, it was not possible to distinguish  
132 between hypersensitivity reactions and infusion reactions in all cases [*see Warnings and*  
133 *Precautions (5.5)*]. Some patients (13%) received premedication, which may have mitigated or

134 masked a hypersensitivity response; however, there is insufficient evidence to determine whether  
135 premedication diminishes the frequency or severity of hypersensitivity reactions.

136

137 BENLYSTA should be administered by healthcare providers prepared to manage anaphylaxis. In  
138 the event of a serious reaction, administration of BENLYSTA must be discontinued immediately  
139 and appropriate medical therapy administered. Patients should be monitored during and for an  
140 appropriate period of time after administration of BENLYSTA. Patients should be informed of  
141 the signs and symptoms of a hypersensitivity reaction and instructed to seek immediate medical  
142 care should a reaction occur.

143

## 144 **5.5 Infusion Reactions**

145 In the controlled clinical trials, adverse events associated with the infusion (occurring on the  
146 same day of the infusion) were reported in 17% (251/1458) of patients receiving BENLYSTA  
147 and 15% (99/675) of patients receiving placebo. Serious infusion reactions (excluding  
148 hypersensitivity reactions) were reported in 0.5% of patients receiving BENLYSTA and 0.4% of  
149 patients receiving placebo and included bradycardia, myalgia, headache, rash, urticaria, and  
150 hypotension. The most common infusion reactions ( $\geq 3\%$  of patients receiving BENLYSTA)  
151 were headache, nausea, and skin reactions. Due to overlap in signs and symptoms, it was not  
152 possible to distinguish between hypersensitivity reactions and infusion reactions in all cases [*see*  
153 *Warnings and Precautions (5.4)*]. Some patients (13%) received premedication, which may have  
154 mitigated or masked an infusion reaction; however there is insufficient evidence to determine  
155 whether premedication diminishes the frequency or severity of infusion reactions [*see Adverse*  
156 *Reactions (6.1)*].

157

158 BENLYSTA should be administered by healthcare providers prepared to manage infusion  
159 reactions. The infusion rate may be slowed or interrupted if the patient develops an infusion  
160 reaction. Healthcare providers should be aware of the risk of hypersensitivity reactions, which  
161 may present as infusion reactions, and monitor patients closely.

162

## 163 **5.6 Depression**

164 In the controlled clinical trials, psychiatric events were reported more frequently with  
165 BENLYSTA (16%) than with placebo (12%), related primarily to depression-related events  
166 (6.3% BENLYSTA and 4.7% placebo), insomnia (6.0% BENLYSTA and 5.3% placebo), and  
167 anxiety (3.9% BENLYSTA and 2.8% placebo). Serious psychiatric events were reported in 0.8%  
168 of patients receiving BENLYSTA (0.6% and 1.2% with 1 and 10 mg/kg, respectively) and 0.4%  
169 of patients receiving placebo. Serious depression was reported in 0.4% (6/1458) of patients  
170 receiving BENLYSTA and 0.1% (1/675) of patients receiving placebo. Two suicides (0.1%)  
171 were reported in patients receiving BENLYSTA. The majority of patients who reported serious  
172 depression or suicidal behavior had a history of depression or other serious psychiatric disorders  
173 and most were receiving psychoactive medications. It is unknown if BENLYSTA treatment is  
174 associated with increased risk for these events.

175

176 Patients receiving BENLYSTA should be instructed to contact their healthcare provider if they  
177 experience new or worsening depression, suicidal thoughts, or other mood changes.

178

179 **5.7 Immunization**

180 Live vaccines should not be given for 30 days before or concurrently with BENLYSTA as  
181 clinical safety has not been established. No data are available on the secondary transmission of  
182 infection from persons receiving live vaccines to patients receiving BENLYSTA or the effect of  
183 BENLYSTA on new immunizations. Because of its mechanism of action, BENLYSTA may  
184 interfere with the response to immunizations.  
185

186 **5.8 Concomitant Use with Other Biologic Therapies or Intravenous  
187 Cyclophosphamide**

188 BENLYSTA has not been studied in combination with other biologic therapies, including B-cell  
189 targeted therapies, or intravenous cyclophosphamide. Therefore, use of BENLYSTA is not  
190 recommended in combination with biologic therapies or intravenous cyclophosphamide.

191 **6 ADVERSE REACTIONS**

192 Because clinical trials are conducted under widely varying conditions, adverse reaction rates  
193 observed in the clinical trials of a drug cannot be directly compared with rates in the clinical  
194 trials of another drug and may not reflect the rates observed in practice.  
195

196 The following have been observed with BENLYSTA and are discussed in detail in the Warnings  
197 and Precautions section:

- 198 • **Mortality** [see Warnings and Precautions (5.1)]
  - 199 • **Serious Infections** [see Warnings and Precautions (5.2)]
  - 200 • **Malignancy** [see Warnings and Precautions (5.3)]
  - 201 • **Hypersensitivity Reactions, Including Anaphylaxis** [see Warnings and Precautions (5.4)]
  - 202 • **Infusion reactions** [see Warnings and Precautions (5.5)]
  - 203 • **Depression** [see Warnings and Precautions (5.6)]
- 204

205 **6.1 Clinical Trials Experience**

206 The data described below reflect exposure to BENLYSTA plus standard of care compared with  
207 placebo plus standard of care in 2133 patients in 3 controlled studies. Patients received  
208 BENLYSTA at doses of 1 mg/kg (N=673), 4 mg/kg (N=111; Trial 1 only), or 10 mg/kg (N=674)  
209 or placebo (N=675) intravenously over a 1-hour period on Days 0, 14, 28, and then every  
210 28 days. In two of the studies (Trial 1 and Trial 3), treatment was given for 48 weeks, while in  
211 the other study (Trial 2) treatment was given for 72 weeks [see Clinical Studies (14)]. Because  
212 there was no apparent dose-related increase in the majority of adverse events observed with  
213 BENLYSTA, the safety data summarized below are presented for the 3 doses pooled, unless  
214 otherwise indicated; the adverse reaction table displays the results for the recommended dose of  
215 10 mg/kg compared with placebo.  
216

217 The population had a mean age of 39 (range 18-75), 94% were female, and 52% were Caucasian.  
218 In these trials, 93% of patients treated with BENLYSTA reported an adverse reaction compared  
219 with 92% treated with placebo.  
220

221 The most common serious adverse reactions were serious infections (6.0% and 5.2% in the  
222 groups receiving BENLYSTA and placebo, respectively) [see Warnings and Precautions (5.2)].  
223

224 The most commonly-reported adverse reactions, occurring in  $\geq 5\%$  of patients in clinical trials  
225 were nausea, diarrhea, pyrexia, nasopharyngitis, bronchitis, insomnia, pain in extremity,  
226 depression, migraine, and pharyngitis.

227  
228 The proportion of patients who discontinued treatment due to any adverse reaction during the  
229 controlled clinical trials was 6.2% for patients receiving BENLYSTA and 7.1% for patients  
230 receiving placebo. The most common adverse reactions resulting in discontinuation of treatment  
231 ( $\geq 1\%$  of patients receiving BENLYSTA or placebo) were infusion reactions (1.6% BENLYSTA  
232 and 0.9% placebo), lupus nephritis (0.7% BENLYSTA and 1.2% placebo), and infections (0.7%  
233 BENLYSTA and 1.0% placebo).

234  
235 Table 1 lists adverse reactions, regardless of causality, occurring in at least 3% of patients with  
236 SLE who received BENLYSTA 10 mg/kg and at an incidence at least 1% greater than that  
237 observed with placebo in the 3 controlled studies.

238 **Table 1. Incidence of Adverse Reactions Occurring in at Least 3% of Patients Treated With BENLYSTA**  
239 **10 mg/kg Plus Standard of Care and at Least 1% More Frequently Than in Patients Receiving Placebo plus**  
240 **Standard of Care in 3 Controlled SLE Studies**

Preferred Term	BENLYSTA 10 mg/kg + Standard of Care (n = 674) %	Placebo + Standard of Care (n = 675) %
Nausea	15	12
Diarrhea	12	9
Pyrexia	10	8
Nasopharyngitis	9	7
Bronchitis	9	5
Insomnia	7	5
Pain in extremity	6	4
Depression	5	4
Migraine	5	4
Pharyngitis	5	3
Cystitis	4	3
Leukopenia	4	2
Gastroenteritis viral	3	1

241

## 242 **6.2 Immunogenicity**

243 In Trials 2 and 3, anti-belimumab antibodies were detected in 4 of 563 (0.7%) patients receiving  
244 BENLYSTA 10 mg/kg and in 27 of 559 (4.8%) patients receiving BENLYSTA 1 mg/kg. The  
245 reported frequency for the group receiving 10 mg/kg may underestimate the actual frequency due  
246 to lower assay sensitivity in the presence of high drug concentrations. Neutralizing antibodies  
247 were detected in 3 patients receiving BENLYSTA 1 mg/kg. Three patients with anti-belimumab  
248 antibodies experienced mild infusion reactions of nausea, erythematous rash, pruritus, eyelid  
249 edema, headache, and dyspnea; none of the reactions was life-threatening. The clinical relevance  
250 of the presence of anti-belimumab antibodies is not known.

251

252 The data reflect the percentage of patients whose test results were positive for antibodies to  
253 belimumab in specific assays. The observed incidence of antibody positivity in an assay is highly  
254 dependent on several factors, including assay sensitivity and specificity, assay methodology,  
255 sample handling, timing of sample collection, concomitant medications, and underlying disease.  
256 For these reasons, comparison of the incidence of antibodies to belimumab with the incidence of  
257 antibodies to other products may be misleading.

258

### 259 **6.3 Postmarketing Experience**

260 The following adverse reactions have been identified during postapproval use of BENLYSTA.  
261 Because these reactions are reported voluntarily from a population of uncertain size, it is not always  
262 possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

- 263 • Fatal anaphylaxis [*see Warnings and Precautions (5.4)*].

## 264 **7 DRUG INTERACTIONS**

265 Formal drug interaction studies have not been performed with BENLYSTA. In clinical trials of  
266 patients with SLE, BENLYSTA was administered concomitantly with other drugs, including  
267 corticosteroids, antimalarials, immunomodulatory and immunosuppressive agents (including  
268 azathioprine, methotrexate, and mycophenolate), angiotensin pathway antihypertensives,  
269 HMG-CoA reductase inhibitors (statins), and NSAIDs without evidence of a clinically  
270 meaningful effect of these concomitant medications on belimumab pharmacokinetics. The effect  
271 of belimumab on the pharmacokinetics of other drugs has not been evaluated [*see*  
272 *Pharmacokinetics (12.3)*].

## 273 **8 USE IN SPECIFIC POPULATIONS**

274

### 275 **8.1 Pregnancy**

276 Pregnancy Category C. There are no adequate and well-controlled clinical studies using  
277 BENLYSTA in pregnant women. Immunoglobulin G (IgG) antibodies, including BENLYSTA,  
278 can cross the placenta. Because animal reproduction studies are not always predictive of human  
279 response, BENLYSTA should be used during pregnancy only if the potential benefit to the  
280 mother justifies the potential risk to the fetus. Women of childbearing potential should use  
281 adequate contraception during treatment with BENLYSTA and for at least 4 months after the  
282 final treatment.

283

284 Nonclinical reproductive studies have been performed in pregnant cynomolgus monkeys  
285 receiving belimumab at doses of 0, 5 and 150 mg/kg by intravenous infusion (the high dose was  
286 approximately 9 times the anticipated maximum human exposure) every 2 weeks from gestation  
287 day 20 to 150. Belimumab was shown to cross the placenta. Belimumab was not associated with  
288 direct or indirect teratogenicity under the conditions tested. Fetal deaths were observed in 14%,  
289 24% and 15% of pregnant females in the 0, 5 and 150 mg/kg groups, respectively. Infant deaths  
290 occurred with an incidence of 0%, 8% and 5%. The cause of fetal and infant deaths is not known.  
291 The relevance of these findings to humans is not known. Other treatment-related findings were  
292 limited to the expected reversible reduction of B cells in both dams and infants and reversible  
293 reduction of IgM in infant monkeys. B-cell numbers recovered after the cessation of belimumab  
294 treatment by about 1 year post-partum in adult monkeys and by 3 months of age in infant  
295 monkeys. IgM levels in infants exposed to belimumab in utero recovered by 6 months of age.

296

297 **Pregnancy Registry:** To monitor maternal-fetal outcomes of pregnant women exposed to  
298 BENLYSTA, a pregnancy registry has been established. Healthcare professionals are encouraged  
299 to register patients and pregnant women are encouraged to enroll themselves by calling  
300 1-877-681-6296.

301

### 302 **8.3 Nursing Mothers**

303 It is not known whether BENLYSTA is excreted in human milk or absorbed systemically after  
304 ingestion. However, belimumab was excreted into the milk of cynomolgus monkeys. Because  
305 maternal antibodies are excreted in human breast milk, a decision should be made whether to  
306 discontinue breastfeeding or to discontinue the drug, taking into account the importance of  
307 breastfeeding to the infant and the importance of the drug to the mother.

308

### 309 **8.4 Pediatric Use**

310 Safety and effectiveness of BENLYSTA have not been established in children.

311

### 312 **8.5 Geriatric Use**

313 Clinical studies of BENLYSTA did not include sufficient numbers of subjects aged 65 or over to  
314 determine whether they respond differently from younger subjects. Use with caution in elderly  
315 patients.

316

### 317 **8.6 Race**

318 In Trial 2 and Trial 3, response rates for the primary endpoint were lower for black subjects in  
319 the BENLYSTA group relative to black subjects in the placebo group [*see Clinical Studies (14)*].  
320 Use with caution in black/African-American patients.

## 321 **10 OVERDOSAGE**

322 There is no clinical experience with overdosage of BENLYSTA. Two doses of up to 20 mg/kg  
323 have been given by intravenous infusion to humans with no increase in incidence or severity of  
324 adverse reactions compared with doses of 1, 4, or 10 mg/kg.

## 325 **11 DESCRIPTION**

326 BENLYSTA (belimumab) is a human IgG1 $\lambda$  monoclonal antibody specific for soluble human B  
327 lymphocyte stimulator protein (BLyS, also referred to as BAFF and TNFSF13B). Belimumab  
328 has a molecular weight of approximately 147 kDa. Belimumab is produced by recombinant DNA  
329 technology in a mammalian cell expression system.

330

331 BENLYSTA is supplied as a sterile, white to off-white, preservative-free, lyophilized powder for  
332 intravenous infusion. Upon reconstitution with Sterile Water for Injection, USP, [*see Dosage  
333 and Administration (2.3)*] each single-use vial delivers 80 mg/mL belimumab in 0.16 mg/mL  
334 citric acid, 0.4 mg/mL polysorbate 80, 2.7 mg/mL sodium citrate, and 80 mg/mL sucrose, with a  
335 pH of 6.5.

336 **12 CLINICAL PHARMACOLOGY**

337

338 **12.1 Mechanism of Action**

339 BENLYSTA is a BLyS-specific inhibitor that blocks the binding of soluble BLyS, a B-cell  
340 survival factor, to its receptors on B cells. BENLYSTA does not bind B cells directly, but by  
341 binding BLyS, BENLYSTA inhibits the survival of B cells, including autoreactive B cells, and  
342 reduces the differentiation of B cells into immunoglobulin-producing plasma cells.

343

344 **12.2 Pharmacodynamics**

345 In Trial 1 and Trial 2 in which B cells were measured, treatment with BENLYSTA significantly  
346 reduced circulating CD19+, CD20+, naïve, and activated B cells, plasmacytoid cells, and the  
347 SLE B-cell subset at Week 52. Reductions in naïve and the SLE B-cell subset were observed as  
348 early as Week 8 and were sustained to Week 52. Memory cells increased initially and slowly  
349 declined toward baseline levels by Week 52. The clinical relevance of these effects on B cells  
350 has not been established.

351

352 Treatment with BENLYSTA led to reductions in IgG and anti-dsDNA, and increases in  
353 complement (C3 and C4). These changes were observed as early as Week 8 and were sustained  
354 through Week 52. The clinical relevance of normalizing these biomarkers has not been  
355 definitively established.

356

357 **12.3 Pharmacokinetics**

358 The pharmacokinetic parameters displayed in Table 2 are based on population parameter  
359 estimates which are specific to the 563 patients who received belimumab 10 mg/kg in Trials 2  
360 and 3 [see *Clinical Studies (14)*].

361 **Table 2. Population Pharmacokinetic Parameters in Patients with SLE after Intravenous Infusion of**  
362 **BENLYSTA 10 mg/kg<sup>1</sup>**

Pharmacokinetic Parameter	Population Estimates (n = 563)
Peak concentration ( $C_{max}$ , $\mu\text{g/mL}$ )	313
Area under the curve ( $AUC_{0-\infty}$ , $\text{day} \bullet \mu\text{g/mL}$ )	3,083
Distribution half-life ( $t_{1/2}$ , days)	1.75
Terminal half-life ( $t_{1/2}$ , days)	19.4
Systemic clearance (CL, mL/day)	215
Volume of distribution ( $V_{ss}$ , L)	5.29

363 <sup>1</sup> Intravenous infusions were administered at 2-week intervals for the first 3 doses and at  
364 4-week intervals thereafter.

365

366 **Drug Interactions:** No formal drug interaction studies have been conducted with belimumab.  
367 Concomitant use of mycophenolate, azathioprine, methotrexate, antimalarials, NSAIDs, aspirin,  
368 and HMG-CoA reductase inhibitors did not significantly influence belimumab pharmacokinetics.  
369 Coadministration of steroids and angiotensin-converting enzyme (ACE) inhibitors resulted in an  
370 increase of systemic clearance of belimumab that was not clinically significant because the  
371 magnitude was well within the range of normal variability of clearance. The effect of belimumab  
372 on the pharmacokinetics of other drugs has not been evaluated.

373

374 **Special Populations:**

375 The following information is based on the population pharmacokinetic analysis.

376

377 *Age:* Age did not significantly influence belimumab pharmacokinetics in the study population,  
378 where the majority of subjects (70%) were between 18 and 45 years of age. No pharmacokinetic  
379 data are available in pediatric patients. Limited pharmacokinetic data are available for elderly  
380 patients as only 1.4% of the subjects included in the pharmacokinetic analysis were 65 years of  
381 age or older [see *Use in Specific Populations (8.5)*].

382

383 *Gender:* Gender did not significantly influence belimumab pharmacokinetics in the largely  
384 (94%) female study population.

385

386 *Race:* Race did not significantly influence belimumab pharmacokinetics. The racial distribution  
387 was 53% white/Caucasian, 16% Asian, 16% Alaska native/American Indian, and 14%  
388 black/African-American.

389

390 *Renal Impairment:* No formal studies were conducted to examine the effects of renal  
391 impairment on the pharmacokinetics of belimumab. Belimumab has been studied in a limited  
392 number of patients with SLE and renal impairment (261 subjects with moderate renal  
393 impairment, creatinine clearance  $\geq 30$  and  $< 60$  mL/min; 14 subjects with severe renal  
394 impairment, creatinine clearance  $\geq 15$  and  $< 30$  mL/min). Although increases in creatinine  
395 clearance and proteinuria ( $> 2$  g/day) increased belimumab clearance, these effects were within  
396 the expected range of variability. Therefore, dosage adjustment in patients with renal impairment  
397 is not recommended.

398

399 *Hepatic Impairment:* No formal studies were conducted to examine the effects of hepatic  
400 impairment on the pharmacokinetics of belimumab. Belimumab has not been studied in patients  
401 with severe hepatic impairment. Baseline ALT and AST levels did not significantly influence  
402 belimumab pharmacokinetics.

403

404 **13 NONCLINICAL TOXICOLOGY**

405

406 **13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility**

407 Long-term animal studies have not been performed to evaluate the carcinogenic potential of  
408 belimumab. The mutagenic potential of belimumab was not evaluated.

409

410 Effects on male and female fertility have not been directly evaluated in animal studies.

411 **14 CLINICAL STUDIES**

412 The safety and effectiveness of BENLYSTA were evaluated in three randomized, double-blind,  
413 placebo-controlled studies involving 2133 patients with SLE according to the American College  
414 of Rheumatology criteria (Trial 1, 2, and 3). Patients with severe active lupus nephritis and  
415 severe active CNS lupus were excluded. Patients were on a stable standard of care SLE treatment  
416 regimen comprising any of the following (alone or in combination): corticosteroids,

417 antimalarials, NSAIDs, and immunosuppressives. Use of other biologics and intravenous  
418 cyclophosphamide were not permitted.

419

420 ***Trial 1: BENLYSTA 1 mg/kg, 4 mg/kg, 10 mg/kg***

421 Trial 1 enrolled 449 patients and evaluated doses of 1, 4, and 10 mg/kg BENLYSTA plus  
422 standard of care compared with placebo plus standard of care over 52 weeks in patients with  
423 SLE. Patients had to have a SELENA-SLEDAI score of  $\geq 4$  at baseline and a history of  
424 autoantibodies (anti-nuclear antibody (ANA) and/or anti-double-stranded DNA (anti-dsDNA),  
425 but 28% of the population was autoantibody negative at baseline. The co-primary endpoints were  
426 percent change in SELENA-SLEDAI score at Week 24 and time to first flare over 52 weeks. No  
427 significant differences between any of the BENLYSTA groups and the placebo group were  
428 observed. Exploratory analysis of this study identified a subgroup of patients (72%), who were  
429 autoantibody positive, in whom BENLYSTA appeared to offer benefit. The results of this study  
430 informed the design of Trials 2 and 3 and led to the selection of a target population and  
431 indication that is limited to autoantibody-positive SLE patients.

432

433 ***Trials 2 and 3: BENLYSTA 1 mg/kg and 10 mg/kg***

434 Trials 2 and 3 were randomized, double-blind, placebo-controlled trials in patients with SLE that  
435 were similar in design except duration - Trial 2 was 76 weeks duration and Trial 3 was 52 weeks  
436 duration. Eligible patients had active SLE disease, defined as a SELENA-SLEDAI score  $\geq 6$ , and  
437 positive autoantibody test results at screening. Patients were excluded from the study if they had  
438 ever received treatment with a B-cell targeted agent or if they were currently receiving other  
439 biologic agents. Intravenous cyclophosphamide was not permitted within the previous 6 months  
440 or during study. Trial 2 was conducted primarily in North America and Europe. Trial 3 was  
441 conducted in South America, Eastern Europe, Asia, and Australia.

442

443 Baseline concomitant medications included corticosteroids (Trial 2: 76%, Trial 3: 96%),  
444 immunosuppressives (Trial 2: 56%, Trial 3: 42%; including azathioprine, methotrexate and  
445 mycophenolate), and antimalarials (Trial 2: 63%, Trial 3: 67%). Most patients (>70%) were  
446 receiving 2 or more classes of SLE medications.

447

448 In Trial 2 and Trial 3, more than 50% of patients had 3 or more active organ systems at baseline.  
449 The most common active organ systems at baseline based on SELENA-SLEDAI were  
450 mucocutaneous (82% in both studies); immunology (Trial 2: 74%, Trial 3: 85%); and  
451 musculoskeletal (Trial 2: 73%, Trial 3: 59%). Less than 16% of patients had some degree of  
452 renal activity and less than 7% of patients had activity in the vascular, cardio-respiratory, or CNS  
453 systems.

454

455 At screening, patients were stratified by disease severity based on their SELENA-SLEDAI score  
456 ( $\leq 9$  vs  $\geq 10$ ), proteinuria level ( $< 2$  g/24 hr vs  $\geq 2$  g/24 hr), and race (African or Indigenous-  
457 American descent vs. other), and then randomly assigned to receive BENLYSTA 1 mg/kg,  
458 BENLYSTA 10 mg/kg, or placebo in addition to standard of care. The patients were  
459 administered study medication intravenously over a 1-hour period on Days 0, 14, 28, and then  
460 every 28 days for 48 weeks in Trial 3 and for 72 weeks in Trial 2.

461

462 The primary efficacy endpoint was a composite endpoint (SLE Responder Index or SRI) that  
463 defined response as meeting each of the following criteria at Week 52 compared with baseline:  
464 •  $\geq 4$ -point reduction in the SELENA-SLEDAI score, and  
465 • no new British Isles Lupus Assessment Group (BILAG) A organ domain score or 2 new  
466 BILAG B organ domain scores, and  
467 • no worsening ( $< 0.30$ -point increase) in Physician's Global Assessment (PGA) score.  
468

469 The SRI uses the SELENA-SLEDAI score as an objective measure of reduction in global disease  
470 activity; the BILAG index to ensure no significant worsening in any specific organ system; and  
471 the PGA to ensure that improvements in disease activity are not accompanied by worsening of  
472 the patient's condition overall.  
473

474 In both Trials 2 and 3, the proportion of SLE patients achieving an SRI response, as defined for  
475 the primary endpoint, was significantly higher in the BENLYSTA 10 mg/kg group than in the  
476 placebo group. The effect on the SRI was not consistently significantly different for the  
477 BENLYSTA 1 mg/kg group relative to placebo in both trials. The 1 mg/kg dose is not  
478 recommended. The trends in comparisons between the treatment groups for the rates of response  
479 for the individual components of the endpoint were generally consistent with that of the SRI  
480 (Table 3). At Week 76 in Trial 2, the SRI response rate with BENLYSTA 10 mg/kg was not  
481 significantly different from that of placebo (39% and 32%, respectively).  
482

483 **Table 3. Clinical Response Rate in Patients with SLE After 52 Weeks of Treatment**

Response <sup>1</sup>	Trial 2			Trial 3		
	Placebo + Standard of Care (n = 275)	BENLYSTA 1 mg/kg + Standard of Care <sup>2</sup> (n = 271)	BENLYSTA 10 mg/kg + Standard of Care (n = 273)	Placebo + Standard of Care (n = 287)	BENLYSTA 1 mg/kg + Standard of Care <sup>2</sup> (n = 288)	BENLYSTA 10 mg/kg + Standard of Care (n = 290)
SLE Responder Index	34%	41%	43%	44%	51%	58%
		(p = 0.104)	(p = 0.021)		(p = 0.013)	(p < 0.001)
Odds Ratio (95% CI) vs. placebo		1.3 (0.9, 1.9)	1.5 (1.1, 2.2)		1.6 (1.1, 2.2)	1.8 (1.3, 2.6)
<b>Components of SLE Responder Index</b>						
Percent of patients with reduction in SELENA-SLEDAI ≥4	36%	43%	47%	46%	53%	58%
Percent of patients with no worsening by BILAG index	65%	75%	69%	73%	79%	81%
Percent of patients with no worsening by PGA	63%	73%	69%	69%	79%	80%

484 <sup>1</sup>Patients dropping out of the study early or experiencing certain increases in background medication were  
485 considered as failures in these analyses. In both studies, a higher proportion of placebo patients were considered as  
486 failures for this reason as compared to the BENLYSTA groups.

487 <sup>2</sup>The 1 mg/kg dose is not recommended.

488  
489 The reduction in disease activity seen in the SRI was related primarily to improvement in the  
490 most commonly involved organ systems namely, mucocutaneous, musculoskeletal, and  
491 immunology.

492  
493 *Effect in Black/African-American Patients:*

494 Exploratory sub-group analyses of SRI response rate in patients of black race were performed. In  
495 Trial 2 and Trial 3 combined, the SRI response rate in black patients (N=148) in the  
496 BENLYSTA groups was less than that in the placebo group (22/50 or 44% for placebo, 15/48 or  
497 31% for BENLYSTA 1 mg/kg, and 18/50 or 36% for BENLYSTA 10 mg/kg). In Trial 1, black  
498 patients (N=106) in the BENLYSTA groups did not appear to have a different response than the  
499 rest of the study population. Although no definitive conclusions can be drawn from these  
500 subgroup analyses, caution should be used when considering BENLYSTA treatment in  
501 black/African-American SLE patients.

502  
503 *Effect on Concomitant Steroid Treatment:*

504 In Trial 2 and Trial 3, 46% and 69% of patients, respectively, were receiving prednisone at doses  
505 > 7.5 mg/day at baseline. The proportion of patients able to reduce their average prednisone dose

506 by at least 25% to  $\leq 7.5$  mg/day during Weeks 40 through 52 was not consistently significantly  
507 different for BENLYSTA relative to placebo in both trials. In Trial 2, 17% of patients receiving  
508 BENLYSTA 10 mg/kg and 19% of patients receiving BENLYSTA 1 mg/kg achieved this level  
509 of steroid reduction compared with 13% of patients receiving placebo. In Trial 3, 19%, 21%, and  
510 12% of patients receiving BENLYSTA 10 mg/kg, BENLYSTA 1 mg/kg, and placebo,  
511 respectively, achieved this level of steroid reduction.

512

#### 513 *Effect on Severe SLE Flares:*

514 The probability of experiencing a severe SLE flare, as defined by a modification of the SELENA  
515 Trial flare criteria which excluded severe flares triggered only by an increase of the SELENA-  
516 SLEDAI score to  $>12$ , was calculated for both Trials 2 and 3. The proportion of patients having  
517 at least 1 severe flare over 52 weeks was not consistently significantly different for BENLYSTA  
518 relative to placebo in both trials. In Trial 2, 18% of patients receiving BENLYSTA 10 mg/kg and  
519 16% of patients receiving BENLYSTA 1 mg/kg had a severe flare compared with 24% of  
520 patients receiving placebo. In Trial 3, 14%, 18%, and 23% of patients receiving BENLYSTA 10  
521 mg/kg, BENLYSTA 1 mg/kg and placebo, respectively, had a severe flare.

## 522 **16 HOW SUPPLIED/STORAGE AND HANDLING**

523 BENLYSTA is a sterile, preservative-free lyophilized powder for reconstitution, dilution, and  
524 intravenous infusion provided in single-use glass vials with a latex-free rubber stopper and a  
525 flip-off seal. Each 5-mL vial contains 120 mg of belimumab. Each 20-mL vial contains 400 mg  
526 of belimumab.

527

528 BENLYSTA is supplied as follows:

120 mg belimumab in a 5-mL single-use vial	NDC 49401-101-01
400 mg belimumab in a 20-mL single-use vial	NDC 49401-102-01

529

530 Store vials of BENLYSTA refrigerated between 2° to 8°C (36° to 46°F). Vials should be  
531 protected from light and stored in the original carton until use. *Do not freeze.* Avoid exposure to  
532 heat. Do not use beyond the expiration date.

## 533 **17 PATIENT COUNSELING INFORMATION**

534 *See FDA-approved patient labeling (Medication Guide)*

535

### 536 **17.1 Advice for the Patient**

537 Patients should be given the Medication Guide for BENLYSTA and provided an opportunity to  
538 read it prior to each treatment session. It is important that the patient's overall health be assessed  
539 at each infusion visit and any questions resulting from the patient's reading of the Medication  
540 Guide be discussed.

541

542 **Mortality:** Patients should be advised that more patients receiving BENLYSTA in the main  
543 clinical trials died than did patients receiving placebo treatment [*see Warnings and Precautions*  
544 (5.1)].

545

546 **Serious Infections:** Patients should be advised that BENLYSTA may decrease their ability to  
547 fight infections. Patients should be asked if they have a history of chronic infections and if they  
548 are currently on any therapy for an infection [*see Warnings and Precautions (5.2)*]. Patients

549 should be instructed to tell their healthcare provider if they develop signs or symptoms of an  
550 infection.

551

552 Hypersensitivity/Anaphylactic and Infusion Reactions: Educate patients on the signs and  
553 symptoms of anaphylaxis, including wheezing, difficulty breathing, peri-oral or lingual edema,  
554 and rash. Patients should be instructed to immediately tell their healthcare provider if they  
555 experience symptoms of an allergic reaction during or after the administration of BENLYSTA  
556 [*see Warnings and Precautions (5.4, 5.5)*].

557

558 Depression: Patients should be instructed to contact their healthcare provider if they experience  
559 new or worsening depression, suicidal thoughts or other mood changes [*see Warnings and*  
560 *Precautions (5.6)*].

561

562 Immunizations: Patients should be informed that they should not receive live vaccines while  
563 taking BENLYSTA. Response to vaccinations could be impaired by BENLYSTA [*see Warnings*  
564 *and Precautions (5.7)*].

565

566 Pregnancy and Nursing Mothers: Patients should be informed that BENLYSTA has not been  
567 studied in pregnant women or nursing mothers so the effects of BENLYSTA on pregnant women  
568 or nursing infants are not known. Patients should be instructed to tell their healthcare provider if  
569 they are pregnant, become pregnant, or are thinking about becoming pregnant [*see Use in*  
570 *Specific Populations (8.1)*]. Patients should be instructed to tell their healthcare provider if they  
571 plan to breastfeed their infant [*see Use in Specific Populations (8.3)*].

572

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574 GlaxoSmithKline.

575

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577 Human Genome Sciences, Inc.  
578 Rockville, Maryland 20850  
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584



GlaxoSmithKline  
Research Triangle Park, NC 27709

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## MEDICATION GUIDE

**BENLYSTA<sup>®</sup>** (ben-LIST-ah)  
**(belimumab)**  
**Injection for intravenous use**

Read this Medication Guide before you start receiving BENLYSTA and before each treatment. There may be new information. This information does not take the place of talking with your healthcare provider about your medical condition or your treatment.

### **What is the most important information I should know about BENLYSTA?**

BENLYSTA can cause serious side effects. Some of these side effects may cause death. It is not known if BENLYSTA causes these serious side effects. Tell your healthcare provider right away if you have any of the symptoms listed below while receiving BENLYSTA.

**1. Infections.** Symptoms of an infection can include:

- fever
- chills
- pain or burning with urination
- urinating often
- bloody diarrhea
- coughing up mucus

**2. Heart Problems.** Symptoms of heart problems can include:

- chest discomfort or pain
- shortness of breath
- cold sweats
- nausea
- dizziness
- discomfort in other areas of the upper body

**3. Mental health problems and suicide.** Symptoms of mental health problems can include:

- thoughts of suicide or dying
- attempt to commit suicide
- trouble sleeping (insomnia)
- new or worse anxiety
- new or worse depression
- acting on dangerous impulses
- other unusual changes in your behavior or mood
- thoughts of hurting yourself or others

## What is BENLYSTA?

BENLYSTA is a prescription medicine used to treat adults with active systemic lupus erythematosus (SLE or lupus) who are receiving other lupus medicines.

BENLYSTA contains belimumab which is in a group of medicines called monoclonal antibodies. Lupus is a disease of the immune system (the body system that fights infection). People with active lupus often have high levels of a certain protein in their blood. BENLYSTA binds to and limits the activity of the protein. When given together with other medicines for lupus, BENLYSTA decreases lupus disease activity more than other lupus medicines alone.

- It is not known if BENLYSTA is safe and effective in people with severe active lupus nephritis or severe active central nervous system lupus.
- It is not known if BENLYSTA is safe and effective in children.

## Who should not receive BENLYSTA?

### Do not receive BENLYSTA if you:

- are allergic to belimumab or any of the ingredients in BENLYSTA. See the end of this Medication Guide for a complete list of ingredients in BENLYSTA.

## What should I tell my healthcare provider before receiving BENLYSTA?

Before you receive BENLYSTA, tell your healthcare provider if you:

- think you have an infection or have infections that keep coming back. You should not receive BENLYSTA if you have an infection unless your healthcare provider tells you to. **See “What is the most important information I should know about BENLYSTA?”**
- have or have had mental health problems such as depression or thoughts of suicide
- have recently received a vaccination or if you think you may need a vaccination. If you are receiving BENLYSTA, you should not receive live vaccines.
- are allergic to other medicines
- are receiving other biologic medicines, monoclonal antibodies or IV infusions of cyclophosphamide (Cytosan®)
- have or have had any type of cancer
- have any other medical conditions
- are pregnant or plan to become pregnant. It is not known if BENLYSTA will harm your unborn baby. Tell your healthcare provider if you become pregnant during your treatment with BENLYSTA.

- If you become pregnant while receiving BENLYSTA, talk to your healthcare provider about enrolling in the BENLYSTA Pregnancy Registry. You can enroll in this registry by calling 1-877-681-6296. The purpose of this registry is to monitor the health of you and your baby.
- are breastfeeding or plan to breastfeed. It is not known if BENLYSTA passes into your breast milk. You and your healthcare provider should decide if you will receive BENLYSTA or breastfeed. You should not do both.

**Tell your healthcare provider about all the medicines you take**, including prescription and non-prescription medicines, vitamins, and herbal supplements.

Know the medicines you take. Keep a list of your medicines with you to show to your healthcare provider and pharmacist when you get a new medicine.

### **How will I receive BENLYSTA?**

- You will be given BENLYSTA by a healthcare provider through a needle placed in a vein (IV infusion). It takes about 1 hour to give you the full dose of BENLYSTA.
- Your healthcare provider will tell you how often you should receive BENLYSTA.
- Your healthcare provider may give you medicines before you receive BENLYSTA to help reduce your chance of having a reaction. A healthcare provider will watch you closely while you are receiving BENLYSTA and after your infusion for signs of a reaction.

### **What are the possible side effects of BENLYSTA?**

**BENLYSTA can cause serious side effects.**

- **See “What is the most important information I should know about BENLYSTA?”**
- **Cancer.** BENLYSTA may reduce the activity of your immune system. Medicines that affect the immune system may increase your risk of certain cancers.
- **Allergic (hypersensitivity) and infusion reactions.** Serious allergic or infusion reactions can happen on the day of or the day after receiving BENLYSTA and may cause death. Tell your healthcare provider right away if you have any of the following symptoms of an allergic or infusion reaction:
  - itching
  - swelling of the face, lips, mouth, tongue, or throat
  - trouble breathing

- anxiousness
- low blood pressure
- dizziness or fainting
- headache
- nausea
- skin rash, redness, or swelling

Your healthcare provider will watch you closely while you are receiving BENLYSTA and after your infusion for signs of a reaction.

**The most common side effects of BENLYSTA include:**

- nausea
- diarrhea
- fever
- stuffy or runny nose
- sore throat
- cough (bronchitis)
- trouble sleeping
- leg or arm pain
- headache (migraine)
- urinary tract infection
- decreased white blood cell count (leukopenia)
- vomiting
- stomach pain

Tell your healthcare provider if you have any side effect that bothers you or that does not go away.

These are not all the possible side effects of BENLYSTA. For more information, ask your healthcare provider.

Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

**General information about the safe and effective use of BENLYSTA**

Medicines are sometimes prescribed for purposes other than those listed in a Medication Guide. Do not use BENLYSTA for a condition for which it was not prescribed.

This Medication Guide summarizes the most important information about BENLYSTA. For more information about BENLYSTA, talk with your healthcare provider.

You can ask your healthcare provider or pharmacist for information about BENLYSTA that is written for healthcare professionals.

For more information about BENLYSTA, go to [www.BENLYSTA.com](http://www.BENLYSTA.com) or call 1-877-423-6597.

### **What are the ingredients in BENLYSTA?**

**Active ingredient:** belimumab.

**Inactive ingredients:** citric acid, polysorbate 80, sodium citrate, sucrose.

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Research Triangle Park, NC 27709

This Medication Guide has been approved by the U.S. Food and Drug Administration.

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