

HIGHLIGHTS OF PRESCRIBING INFORMATION

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

Privigen, Immune Globulin Intravenous (Human), 10% Liquid
Initial U.S. Approval: 2007

WARNING: ACUTE RENAL DYSFUNCTION/FAILURE

See full prescribing information for complete boxed warning.

- Renal dysfunction, acute renal failure, osmotic nephropathy, and death may occur with the administration of human immune globulin intravenous (IGIV) products.
- Renal dysfunction and acute renal failure occur more commonly in patients receiving IGIV products that contain sucrose. Privigen does not contain sucrose.
- For patients at risk of renal dysfunction or renal failure, administer Privigen at the minimum infusion rate practicable.

RECENT MAJOR CHANGES

Warnings and Precautions (5.6, 5.8) 05/2012

INDICATIONS AND USAGE

Privigen is an Immune Globulin Intravenous (Human), 10% Liquid indicated for the treatment of:

- Primary humoral immunodeficiency (PI) (1.1)
- Chronic immune thrombocytopenic purpura (ITP) (1.2)

DOSAGE AND ADMINISTRATION

Intravenous Use Only

Indication	Dose (2.2)	Initial Infusion Rate (2.3)	Maintenance Infusion Rate (if tolerated) (2.3)
PI	200-800 mg/kg (2-8 mL/kg) every 3-4 weeks	0.5 mg/kg/min (0.005 mL/kg/min)	Increase to 8 mg/kg/min (0.08 mL/kg/min)
ITP	1 g/kg (10 mL/kg) for 2 consecutive days	0.5 mg/kg/min (0.005 mL/kg/min)	Increase to 4 mg/kg/min (0.04 mL/kg/min)

- Ensure that patients with pre-existing renal insufficiency are not volume depleted, and discontinue Privigen if renal function deteriorates (2.3, 5.2).
- For patients at risk of renal dysfunction or thrombotic events, administer Privigen at the minimum infusion rate practicable (2.3, 5.2, 5.4).

DOSAGE FORMS AND STRENGTHS

Privigen is a liquid solution containing 10% IgG (0.1 g/mL) (3).

CONTRAINDICATIONS

- History of anaphylactic or severe systemic reaction to human immune globulin (4)
- Hyperprolinemia (Privigen contains the stabilizer L-proline) (4)
- IgA-deficient patients with antibodies to IgA and a history of hypersensitivity (4)

WARNINGS AND PRECAUTIONS

- IgA-deficient patients with antibodies to IgA are at greater risk of developing severe hypersensitivity and anaphylactic reactions (5.1).
- Monitor renal function, including blood urea nitrogen and serum creatinine, and urine output in patients at risk of developing acute renal failure (5.2).
- Thrombotic events may occur. Monitor patients with known risk factors for thrombotic events; consider baseline assessment of blood viscosity for those at risk of hyperviscosity (5.3).
- Hyperproteinemia, increased serum viscosity, and hyponatremia may occur (5.4).
- Aseptic meningitis syndrome (AMS) may occur, especially with high doses or rapid infusion (5.5).
- Hemolysis that is either intravascular or due to enhanced red blood cell sequestration can develop subsequent to Privigen treatments. Risk factors for hemolysis include high doses and non-O blood group. Closely monitor patients for hemolysis and hemolytic anemia (5.6).
- Monitor patients for pulmonary adverse reactions (transfusion-related acute lung injury [TRALI]) (5.7).
- Carefully consider the relative risks and benefits before prescribing the high dose regimen (for chronic ITP) in patients at increased risk of thrombosis, hemolysis, acute kidney injury, or volume overload (5.8).
- Privigen is made from human blood and may contain infectious agents, e.g., viruses and, theoretically, the Creutzfeldt-Jakob disease (CJD) agent (5.9).

ADVERSE REACTIONS

- PI** – The most common adverse reactions, observed in >5% of study subjects, were headache, pain, nausea, fatigue, chills, vomiting, joint swelling/effusion, pyrexia, and urticaria. Serious adverse reactions were hypersensitivity, chills, fatigue, dizziness, and increased body temperature (6).
- Chronic ITP** – The most common adverse reactions, observed in >5% of study subjects, were headache, pyrexia/hyperthermia, positive direct antiglobulin test (DAT), anemia, vomiting, nausea, bilirubin conjugated increased, bilirubin unconjugated increased, hyperbilirubinemia, and blood lactate dehydrogenase increased. A serious adverse reaction was aseptic meningitis (6).

To report SUSPECTED ADVERSE REACTIONS, contact CSL Behring Pharmacovigilance at 1-866-915-6958 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

DRUG INTERACTIONS

The passive transfer of antibodies may:

- Lead to misinterpretation of the results of serological testing (5.10, 7.2).
- Interfere with the response to live virus vaccines (7.1).

USE IN SPECIFIC POPULATIONS

- Pregnancy:** No human or animal data. Use only if clearly needed (8.1).
- In patients over age 65 or in any patient at risk of developing renal insufficiency, do not exceed the recommended dose, and infuse Privigen at the minimum rate practicable (8.5).

See 17 for PATIENT COUNSELING INFORMATION.

Revised: May 2012

FULL PRESCRIBING INFORMATION: CONTENTS*

WARNING – ACUTE RENAL DYSFUNCTION/FAILURE

1 INDICATIONS AND USAGE

- Primary Humoral Immunodeficiency
- Chronic Immune Thrombocytopenic Purpura

2 DOSAGE AND ADMINISTRATION

- Preparation and Handling
- Dosage
- Administration

3 DOSAGE FORMS AND STRENGTHS

4 CONTRAINDICATIONS

5 WARNINGS AND PRECAUTIONS

- Hypersensitivity
- Renal Dysfunction/Failure
- Thrombotic Events
- Hyperproteinemia, Increased Serum Viscosity, and Hyponatremia
- Aseptic Meningitis Syndrome (AMS)
- Hemolysis
- Transfusion-Related Acute Lung Injury (TRALI)
- Volume Overload
- Transmissible Infectious Agents
- Interference with Laboratory Tests

6 ADVERSE REACTIONS

- Clinical Trials Experience
- Postmarketing Experience

7 DRUG INTERACTIONS

- Live Virus Vaccines
- Serological Testing

8 USE IN SPECIFIC POPULATIONS

- Pregnancy
- Nursing Mothers
- Pediatric Use
- Geriatric Use

10 OVERDOSAGE

11 DESCRIPTION

12 CLINICAL PHARMACOLOGY

- Mechanism of Action
- Pharmacokinetics

14 CLINICAL STUDIES

- Treatment of Primary Humoral Immunodeficiency
- Treatment of Chronic Immune Thrombocytopenic Purpura

15 REFERENCES

16 HOW SUPPLIED/STORAGE AND HANDLING

17 PATIENT COUNSELING INFORMATION

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

FULL PRESCRIBING INFORMATION

WARNING: ACUTE RENAL DYSFUNCTION/FAILURE

- Use of Immune Globulin Intravenous (IGIV) products, particularly those containing sucrose, have been reported to be associated with renal dysfunction, acute renal failure, osmotic nephropathy, and death.¹ Patients at risk of acute renal failure include those with any degree of pre-existing renal insufficiency, diabetes mellitus, advanced age (above 65 years of age), volume depletion, sepsis, paraproteinemia, or receiving known nephrotoxic drugs (*see Warnings and Precautions [5.2]*). Privigen does not contain sucrose.
- For patients at risk of renal dysfunction or failure, administer Privigen at the minimum infusion rate practicable (*see Dosage and Administration [2.3], Warnings and Precautions [5.2]*).

1 INDICATIONS AND USAGE

Privigen is an Immune Globulin Intravenous (Human), 10% Liquid indicated for the treatment of the following conditions.

1.1 Primary Humoral Immunodeficiency

Privigen is indicated as replacement therapy for primary humoral immunodeficiency (PI). This includes, but is not limited to, the humoral immune defect in congenital agammaglobulinemia, common variable immunodeficiency (CVID), X-linked agammaglobulinemia, Wiskott-Aldrich syndrome, and severe combined immunodeficiencies.

1.2 Chronic Immune Thrombocytopenic Purpura

Privigen is indicated for the treatment of patients with chronic immune thrombocytopenic purpura (ITP) to raise platelet counts.

2 DOSAGE AND ADMINISTRATION

2.1 Preparation and Handling

- Privigen is a clear or slightly opalescent, colorless to pale yellow solution. Inspect parenteral drug products visually for particulate matter and discoloration prior to administration, whenever solution and container permit. Do not use if the solution is cloudy, turbid, or if it contains particulate matter.
- DO NOT SHAKE.
- Do not freeze. Do not use if Privigen has been frozen.
- Privigen should be at room temperature (up to 25°C [77°F]) at the time of administration.
- Do not use Privigen beyond the expiration date on the product label.

- 34 • The Privigen vial is for single-use only. Promptly use any vial that
35 has been entered. Privigen contains no preservative. Discard
36 partially used vials or unused product in accordance with local
37 requirements.
- 38 • Infuse Privigen using a separate infusion line. Prior to use, the
39 infusion line may be flushed with Dextrose Injection, USP (D5W) or
40 0.9% Sodium Chloride for Injection, USP.
- 41 • Do not mix Privigen with other IGIV products or other intravenous
42 medications. However, Privigen may be diluted with Dextrose
43 Injection, USP (D5W).
- 44 • An infusion pump may be used to control the rate of administration.
- 45 • If large doses of Privigen are to be administered, several vials may
46 be pooled using aseptic technique. Begin infusion within 8 hours of
47 pooling.

48

49 **2.2 Dosage**

50 **Treatment of Primary Humoral Immunodeficiency (PI)**

51 As there are significant differences in the half-life of IgG among patients
52 with PI, the frequency and amount of immunoglobulin therapy may vary
53 from patient to patient. The proper amount can be determined by
54 monitoring clinical response.

55

56 The recommended dose of Privigen for patients with PI is 200 to 800
57 mg/kg (2 to 8 mL/kg), administered every 3 to 4 weeks. If a patient
58 misses a dose, administer the missed dose as soon as possible, and then
59 resume scheduled treatments every 3 or 4 weeks, as applicable.

60

61 Adjust the dosage over time to achieve the desired serum IgG trough
62 levels and clinical responses. No randomized, controlled trial data are
63 available to determine an optimal trough level in patients receiving
64 immune globulin therapy.

65

66 **Treatment of Chronic Immune Thrombocytopenic Purpura (ITP)**

67 The recommended dose of Privigen for patients with chronic ITP is 1 g/kg
68 (10 mL/kg) administered daily for 2 consecutive days, resulting in a total
69 dosage of 2 g/kg.

70

71 Carefully consider the relative risks and benefits before prescribing the
72 high dose regimen (e.g., 1 g/kg/day for 2 days) in patients at increased risk
73 of thrombosis, hemolysis, acute kidney injury, or volume overload (see
74 *Warnings and Precautions [5.8]*).

75

76

77

78 **2.3 Administration**79 **Privigen is for intravenous administration only.**

80 Monitor the patient's vital signs throughout the infusion. Slow or stop the
81 infusion if adverse reactions occur. If symptoms subside promptly, the
82 infusion may be resumed at a lower rate that is comfortable for the patient.

83

84 Ensure that patients with pre-existing renal insufficiency are not volume
85 depleted. For patients judged to be at risk for renal dysfunction or
86 thrombotic events, administer Privigen at the minimum infusion rate
87 practicable, and discontinue Privigen administration if renal function
88 deteriorates (see *Boxed Warning, Warnings and Precautions [5.2, 5.3]*).

89

90 Table 1 provides the recommended infusion rates for Privigen.

91

92 **Table 1: Recommended Infusion Rates for Privigen**

93

Indication	Dose	Initial infusion rate	Maintenance infusion rate (if tolerated)
PI	200-800 mg/kg (2-8 mL/kg) every 3-4 weeks	0.5 mg/kg/min (0.005 mL/kg/min)	Increase to 8 mg/kg/min (0.08 mL/kg/min)
ITP	1 g/kg (10 mL/kg) for 2 consecutive days	0.5 mg/kg/min (0.005 mL/kg/min)	Increase to 4 mg/kg/min (0.04 mL/kg/min)

94

95 The following patients may be at risk of developing systemic reactions
96 (mimicking symptoms of an inflammatory response or infection) on rapid
97 infusion of Privigen (greater than 4 mg/kg/min [0.04 mL/kg/min]): 1)
98 those who have never received Privigen or another IgG product or who
99 have not received it within the past 8 weeks, and 2) those who are
100 switching from another IgG product. These patients should be started at a
101 slow rate of infusion (e.g., 0.5 mg/kg/min [0.005 mL/kg/min] or less) and
102 gradually advanced to the maximum rate as tolerated.

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105 **3 DOSAGE FORMS AND STRENGTHS**

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107 Privigen is a liquid solution containing 10% IgG (0.1 g/mL) for
108 intravenous infusion.

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4 CONTRAINDICATIONS

- Privigen is contraindicated in patients who have a history of anaphylactic or severe systemic reaction to the administration of human immune globulin.
- Privigen is contraindicated in patients with hyperprolinemia because it contains the stabilizer L-proline (see *Description [11]*).
- Privigen is contraindicated in IgA-deficient patients with antibodies to IgA and a history of hypersensitivity (see *Warnings and Precautions [5.1]*).

5 WARNINGS AND PRECAUTIONS

5.1 Hypersensitivity

Severe hypersensitivity reactions may occur (see *Contraindications [4]*). In case of hypersensitivity, discontinue the Privigen infusion immediately and institute appropriate treatment. Medications such as epinephrine should be available for immediate treatment of acute hypersensitivity reactions.

Privigen contains trace amounts of IgA (≤ 25 mcg/mL) (see *Description [11]*). Individuals with IgA deficiency can develop anti-IgA antibodies and anaphylactic reactions (including anaphylaxis and shock) after administration of blood components containing IgA. Patients with known antibodies to IgA may have a greater risk of developing potentially severe hypersensitivity and anaphylactic reactions with administration of Privigen. Privigen is contraindicated in patients with antibodies against IgA and a history of hypersensitivity.

5.2 Renal Dysfunction/Failure

Acute renal dysfunction/failure, osmotic nephropathy, and death may occur with the use of IGIV products, including Privigen, and particularly in those products containing sucrose. (Privigen does not contain sucrose.) Ensure that patients are not volume depleted and assess renal function, including measurement of blood urea nitrogen (BUN) and serum creatinine, before the initial infusion of Privigen and at appropriate intervals thereafter.

Periodic monitoring of renal function and urine output is particularly important in patients judged to be at increased risk of developing acute renal failure.¹ If renal function deteriorates, consider discontinuing Privigen. For patients judged to be at risk of developing renal dysfunction because of pre-existing renal insufficiency, or predisposition to acute renal

156 failure (such as those with diabetes mellitus or hypovolemia, those who
157 are obese, those who use concomitant nephrotoxic medicinal products, or
158 those who are over 65 years of age), administer Privigen at the minimum
159 rate of infusion practicable (*see [Boxed Warning](#), [Dosage and](#)*
160 *[Administration \[2.3\]](#)*).

161

162 **5.3 Thrombotic Events**

163 Thrombotic events may occur following treatment with IGIV products,
164 including Privigen.²⁻⁴ Patients at risk include those with a history of
165 atherosclerosis, multiple cardiovascular risk factors, advanced age,
166 impaired cardiac output, coagulation disorders, prolonged periods of
167 immobilization, and/or known/suspected hyperviscosity.

168

169 Because of the potentially increased risk of thrombosis, consider baseline
170 assessment of blood viscosity in patients at risk for hyperviscosity,
171 including those with cryoglobulins, fasting chylomicronemia/markedly
172 high triacylglycerols (triglycerides), or monoclonal gammopathies. For
173 patients judged to be at risk of developing thrombotic events, administer
174 Privigen at the minimum rate of infusion practicable (*see [Dosage and](#)*
175 *[Administration \[2.3\]](#)*).

176

177 **5.4 Hyperproteinemia, Increased Serum Viscosity, and** 178 **Hyponatremia**

179 Hyperproteinemia, increased serum viscosity, and hyponatremia may
180 occur following treatment with IGIV products, including Privigen. The
181 hyponatremia is likely to be a pseudohyponatremia, as demonstrated by a
182 decreased calculated serum osmolality or elevated osmolar gap. It is
183 critical to distinguish true hyponatremia from pseudohyponatremia, as
184 treatment aimed at decreasing serum free water in patients with
185 pseudohyponatremia may lead to volume depletion, a further increase in
186 serum viscosity, and a possible predisposition to thromboembolic events.⁵

187

188 **5.5 Aseptic Meningitis Syndrome (AMS)**

189 AMS may occur infrequently following treatment with Privigen (*see*
190 *[Adverse Reactions \[6\]](#)*) and other human immune globulin products.
191 Discontinuation of treatment has resulted in remission of AMS within
192 several days without sequelae.⁶ AMS usually begins within several hours
193 to 2 days following IGIV treatment.

194

195 AMS is characterized by the following signs and symptoms: severe
196 headache, nuchal rigidity, drowsiness, fever, photophobia, painful eye
197 movements, nausea, and vomiting. Cerebrospinal fluid (CSF) studies are
198 frequently positive with pleocytosis up to several thousand cells per cubic
199 millimeter, predominantly from the granulocytic series, and with elevated
200 protein levels up to several hundred mg/dL, but negative culture results.

201 Conduct a thorough neurological examination on patients exhibiting such
202 signs and symptoms, including CSF studies, to rule out other causes of
203 meningitis.

204
205 AMS may occur more frequently in association with high doses (2 g/kg)
206 and/or rapid infusion of IGIV.

207 208 **5.6 Hemolysis**

209 Privigen may contain blood group antibodies that can act as hemolysins
210 and induce *in vivo* coating of red blood cells (RBCs) with
211 immunoglobulin, causing a positive direct antiglobulin test (DAT)
212 (Coombs' test) result and hemolysis.⁷⁻⁹ Delayed hemolytic anemia can
213 develop subsequent to Privigen therapy due to enhanced RBC
214 sequestration, and acute hemolysis, consistent with intravascular
215 hemolysis, has been reported.¹⁰ Cases of severe hemolysis-related renal
216 dysfunction/failure or disseminated intravascular coagulation have
217 occurred following infusion of Privigen.

218
219 The following risk factors may be associated with the development of
220 hemolysis: high doses (e.g., ≥ 2 g/kg), given either as a single
221 administration or divided over several days, and non-O blood group.¹¹
222 Other individual patient factors, such as an underlying inflammatory state
223 (as may be reflected by, for example, elevated C-reactive protein or
224 erythrocyte sedimentation rate), have been hypothesized to increase the
225 risk of hemolysis following administration of IGIV,¹² but their role is
226 uncertain. Hemolysis has been reported following administration of IGIV
227 for a variety of indications, including ITP and PI.⁹

228
229 Closely monitor patients for clinical signs and symptoms of hemolysis,
230 particularly patients with risk factors noted above. Consider appropriate
231 laboratory testing in higher risk patients, including measurement of
232 hemoglobin or hematocrit prior to infusion and within approximately 36 to
233 96 hours post infusion. If clinical signs and symptoms of hemolysis or a
234 significant drop in hemoglobin or hematocrit have been observed, perform
235 additional confirmatory laboratory testing. If transfusion is indicated for
236 patients who develop hemolysis with clinically compromising anemia
237 after receiving IGIV, perform adequate cross-matching to avoid
238 exacerbating on-going hemolysis.

239 240 **5.7 Transfusion-Related Acute Lung Injury (TRALI)**

241 Noncardiogenic pulmonary edema may occur following treatment with
242 IGIV products, including Privigen.¹³ TRALI is characterized by severe
243 respiratory distress, pulmonary edema, hypoxemia, normal left ventricular
244 function, and fever. Symptoms typically appear within 1 to 6 hours
245 following treatment.

246

247 Monitor patients for pulmonary adverse reactions. If TRALI is suspected,
248 perform appropriate tests for the presence of anti-neutrophil antibodies
249 and anti-human leukocyte antigen (HLA) antibodies in both the product
250 and the patient's serum.

251

252 TRALI may be managed using oxygen therapy with adequate ventilatory
253 support.

254

255 **5.8 Volume Overload**

256 Carefully consider the relative risks and benefits before prescribing the
257 high dose regimen (for chronic ITP) in patients at increased risk of
258 thrombosis, hemolysis, acute kidney injury, or volume overload.

259

260 **5.9 Transmissible Infectious Agents**

261 Because Privigen is made from human blood, it may carry a risk of
262 transmitting infectious agents (e.g., viruses and, theoretically, the
263 Creutzfeldt-Jakob disease [CJD] agent). The risk of infectious agent
264 transmission has been reduced by screening plasma donors for prior
265 exposure to certain viruses, testing for the presence of certain current virus
266 infections, and including virus inactivation/removal steps in the
267 manufacturing process for Privigen.

268

269 Report any infection thought to be possibly transmitted by Privigen to
270 CSL Behring Pharmacovigilance at 1-866-915-6958.

271

272 **5.10 Interference with Laboratory Tests**

273 Various passively transferred antibodies in immunoglobulin preparations
274 may lead to misinterpretation of the results of serological testing.

275

276

277 **6 ADVERSE REACTIONS**

278

279 The most serious adverse reaction observed in clinical study subjects
280 receiving Privigen for PI was hypersensitivity in one subject. The most
281 common adverse reactions observed in >5% of clinical study subjects with
282 PI were headache, pain, nausea, fatigue, chills, vomiting, joint
283 swelling/effusion, pyrexia, and urticaria.

284

285 The most serious adverse reactions observed in clinical study subjects
286 receiving Privigen for chronic ITP were aseptic meningitis syndrome in
287 one subject and hemolysis in two subjects. A total of 8 subjects (14%) in
288 the ITP study experienced hemolysis as documented from clinical
289 laboratory data. The most common adverse reactions observed in >5% of
290 clinical study subjects with chronic ITP were headache,

291 pyrexia/hyperthermia, positive DAT, anemia, vomiting, nausea,
292 hyperthermia, bilirubin conjugated increased, bilirubin unconjugated
293 increased, hyperbilirubinemia, and blood lactate dehydrogenase increased.
294

295 **6.1 Clinical Trials Experience**

296 *Because different clinical trials are conducted under widely varying*
297 *conditions, adverse reaction rates observed in the clinical trials of a drug*
298 *cannot be directly compared to rates in the clinical trials of another drug*
299 *and may not reflect the rates observed in clinical practice.*
300

301 Treatment of Primary Humoral Immunodeficiency

302 In a prospective, open-label, single-arm, multicenter clinical study (pivotal
303 study), 80 subjects with PI (with a diagnosis of XLA or CVID) received
304 Privigen every 3 or 4 weeks for up to 12 months (*see Clinical Studies*
305 *[14.1]*). All subjects had been on regular IGIV replacement therapy for at
306 least 6 months prior to participating in the study. Subjects ranged in age
307 from 3 to 69; 46 (57.5%) were male and 34 (42.5%) were female.
308

309 The safety analysis included all 80 subjects, 16 (20%) on the 3-week
310 schedule and 64 (80%) on the 4-week schedule. The median dose of
311 Privigen administered was 428.3 mg/kg (3-week schedule) or 440.6 mg/kg
312 (4-week schedule) and ranged from 200 to 888 mg/kg. A total of 1038
313 infusions of Privigen were administered, 272 in the 3-week schedule and
314 766 in the 4-week schedule.
315

316 Routine premedication was not allowed. However, subjects who
317 experienced two consecutive infusion-related adverse events (AEs) that
318 were likely to be prevented by premedication were permitted to receive
319 antipyretics, antihistamines, NSAIDs, or antiemetic agents. During the
320 study, 8 (10%) subjects received premedication prior to 51 (4.9%) of the
321 1038 infusions administered.
322

323 Temporally associated AEs are those occurring during an infusion or
324 within 72 hours after the end of an infusion, irrespective of causality. In
325 this study, the upper bound of the 1-sided 97.5% confidence interval for
326 the proportion of Privigen infusions temporally associated with one or
327 more AEs was 23.8% (actual proportion: 20.8%). The total number of
328 temporally associated AEs was 397 (a rate of 0.38 AEs per infusion),
329 reflecting that some subjects experienced more than one AE during the
330 observation period.
331

332

333 Table 2 lists the temporally associated AEs that occurred in >5% of
334 subjects, *irrespective of causality*.

335

336 **Table 2: PI Pivotal Study – Adverse Events* Occurring in >5% of**
337 **Subjects During a Privigen Infusion or Within 72 Hours**
338 **After the End of an Infusion, *Irrespective of Causality***

339

Adverse Event	Number (%) of Subjects [n=80]	Number (Rate) of Infusions with Adverse Event [n=1038]
Headache	35 (43.8)	82 (0.079)
Pain	20 (25.0)	44 (0.042)
Fatigue	13 (16.3)	27 (0.026)
Nausea	10 (12.5)	19 (0.018)
Chills	9 (11.3)	15 (0.014)
Vomiting	7 (8.8)	13 (0.013)
Pyrexia	6 (7.5)	10 (0.010)
Cough	5 (6.3)	5 (0.005)
Diarrhea	5 (6.3)	5 (0.005)
Stomach discomfort	5 (6.3)	5 (0.005)

340

* Excluding infections.

341

342 Of the 397 temporally associated AEs reported for the 80 subjects with PI,
343 the investigators judged 192 to be at least possibly related to the infusion
344 of Privigen (including 5 serious, severe AEs described below). Of these,
345 91 were mild, 81 were moderate, 19 were severe, and 1 was of unknown
346 severity.

347

348 Table 3 lists the adverse reactions (AEs considered to be “at least possibly
349 related” to the infusion of Privigen) that occurred in >5% of subjects with
350 PI, *irrespective of time of occurrence*.

351

352 **Table 3: PI Pivotal Study – Adverse Reactions Considered “at Least**
353 **Possibly Related” to Privigen Occurring in >5% of Subjects,**
354 ***Irrespective of Time of Occurrence***

355

Adverse Reaction	Number (%) of Subjects [n=80]	Number (Rate) of Infusions with Adverse Reaction [n=1038]
Headache	24 (30.0)	62 (0.060)
Pain, all types*	12 (15.0) [†]	26 (0.025)
Nausea	10 (12.5)	18 (0.017)
Fatigue	9 (11.3)	16 (0.015)
Chills	9 (11.3)	15 (0.014)
Vomiting	6 (7.5)	11 (0.011)

356

* Includes abdominal pain lower, abdominal tenderness, arthralgia, back pain, chest pain, infusion-site pain, injection-site pain, neck pain, pain, pain in extremity, and pharyngolaryngeal pain.

357

358 † Some subjects experienced more than one type of pain.

359

360 Sixteen (20%) subjects experienced 41 serious AEs (SAEs). Five of these
361 SAEs (hypersensitivity, chills, fatigue, dizziness, and increased body
362 temperature, all severe) were related to Privigen, occurred in one subject,
363 and resulted in the subject's withdrawal from the study. Two other
364 subjects withdrew from the study due to AEs related to Privigen treatment
365 (chills and headache in one subject; vomiting in the other).

366

367 Seventy-seven of the 80 subjects enrolled in this study had a negative
368 DAT at baseline. Of these 77 subjects, 36 (46.8%) developed a positive
369 DAT at some time during the study. However, no subjects showed
370 evidence of hemolytic anemia.

371

372 During this study, no subjects tested positive for infection due to human
373 immunodeficiency virus (HIV), hepatitis B virus (HBV), hepatitis C virus
374 (HCV), or B19 virus (B19V).

375

376 An extension of the pivotal study was conducted in 55 adult and pediatric
377 subjects with PI to collect additional efficacy, safety, and tolerability data.
378 This study included 45 subjects from the pivotal study who were receiving
379 Privigen and 10 new subjects who were receiving another IGIV product
380 prior to enrolling in the extension study. Subjects ranged in age from 4 to
381 81 years; 26 (47.3%) were male and 29 (52.7%) were female.

382

383 Subjects were treated with Privigen at median doses ranging from 286 to
384 832 mg/kg per infusion over a treatment period ranging from 1 to 27
385 months. Twelve (21.8%) subjects were on a 3-week treatment schedule
386 with the number of infusions per subject ranging from 4 to 38 (median: 8
387 infusions); 43 (78.2%) subjects were on a 4-week schedule with the
388 number of infusions ranging from 1 to 31 (median: 15 infusions). A total
389 of 771 infusions were administered in this study.

390

391 In this study, subjects who continued from the pivotal study were
392 permitted to receive infusions of Privigen at a rate up to 12 mg/kg/min (as
393 opposed to the maximum of 8 mg/kg/min allowed in the pivotal study) at
394 the discretion of the investigator based on individual tolerability. Twenty-
395 three (51%) of the 45 subjects from the pivotal study (41.8% of the 55
396 subjects in the extension study) received 265 (38.4%) infusions at a
397 maximum rate greater than the recommended rate of 8 mg/kg/min (*see*
398 *Dosing and Administration [2.3]*). The median of the maximum infusion
399 rate in this subset was 12 mg/kg/min. However, because the study was not
400 designed to compare infusion rates, no definitive conclusions regarding
401 tolerability could be drawn for infusion rates higher than the
402 recommended rate of 8 mg/kg/min.

403

404 In this study, the proportion of infusions temporally associated with one or
405 more AEs occurring during a Privigen infusion or within 72 hours after the
406 end of an infusion was 15%. The total number of temporally associated
407 AEs, irrespective of causality, was 206 (a rate of 0.27 AEs per infusion),
408 reflecting that some subjects experienced more than one AE during the
409 observation period.

410

411 Table 4 lists the temporally associated AEs that occurred in >5% of
412 subjects, *irrespective of causality*.

413

414 **Table 4: PI Extension Study – Adverse Events* Occurring in >5% of**
415 **Subjects During a Privigen Infusion or Within 72 Hours After**
416 **the End of an Infusion, *Irrespective of Causality***

417

Adverse Event*	Number (%) of Subjects [n=55]	Number (Rate) of Infusions with Adverse Event [n=771]
Headache	18 (32.7)	56 (0.073)
Pain, all types [†]	14 (25.5) [‡]	31 (0.040)
Abdominal pain [§]	3 (5.5)	4 (0.005)
Chest pain	3 (5.5)	4 (0.005)
Pharyngolaryngeal pain	3 (5.5)	4 (0.005)
Nausea	6 (10.9)	10 (0.013)
Pyrexia	4 (7.3)	9 (0.012)
Chills	3 (5.5)	7 (0.009)
Influenza-like illness	3 (5.5)	4 (0.005)

418 Note: The AE rates in this study cannot be compared directly to the rates in other IGIV studies,
419 including the original pivotal study described earlier in this section, because (1) the extension study
420 used an enriched population and (2) the selective use of higher infusion rates at the investigators'
421 discretion in a subset of subjects may have introduced bias.

422 * Excluding infections.

423 † Includes abdominal pain, abdominal pain upper, arthralgia, back pain, chest pain, fibromyalgia,
424 injection-site pain, myalgia, pain, pain in extremity, painful respiration, pharyngolaryngeal pain, and
425 toothache.

426 ‡ Some subjects experienced more than one type of pain.

427 § Also includes abdominal pain, upper.

428

429 Of the 206 temporally associated AEs reported for the 55 subjects with PI,
430 the investigators judged 125 to be at least possibly related to the infusion
431 of Privigen. Of these, 76 were mild, 40 were moderate, and 9 were severe.

432

433
434 Table 5 lists the adverse reactions (AEs considered to be “at least possibly
435 related” to the infusion of Privigen) that occurred in >5% of subjects,
436 *irrespective of time of occurrence.*

437
438 **Table 5: PI Extension Study – Adverse Reactions Considered “at
439 Least Possibly Related” to Privigen Occurring in >5% of
440 Subjects, Irrespective of Time of Occurrence**

441

Adverse Reaction	Number (%) of Subjects [n=55]	Number (Rate) of Infusions With Adverse Reaction [n=771]
Headache	16 (29.1)	53 (0.069)
Pain, all types*	11 (20.0) [†]	26 (0.034)
Abdominal pain [‡]	4 (7.3)	6 (0.008)
Chest pain	3 (5.5)	4 (0.005)
Chills	3 (5.5)	7 (0.009)
Fatigue	3 (5.5)	5 (0.006)
Joint swelling/effusion	3 (5.5)	7 (0.009)
Pyrexia	3 (5.5)	10 (0.013)
Urticaria	3 (5.5)	4 (0.005)

442 * Includes abdominal pain, abdominal pain lower, abdominal pain upper, arthralgia, back pain, chest
443 pain, injection-site pain, musculoskeletal pain, myalgia, pain, and painful respiration.

444 [†] Some subjects experienced more than one type of pain.

445 [‡] Includes abdominal pain, lower and abdominal pain, upper.

446

447 Eleven (20%) subjects experienced 17 SAEs, none of which were
448 considered to be related to Privigen. Three subjects experienced AEs that
449 were considered to be at least possibly related to Privigen: dyspnea and
450 pancytopenia in one subject, a transient ischemic attack 16 days after the
451 infusion in one subject, and mild urticaria in one subject, resulting in the
452 subject’s withdrawal from the study.

453

454 Treatment of Chronic Immune Thrombocytopenic Purpura

455 In a prospective, open-label, single-arm, multicenter clinical study, 57
456 subjects with chronic ITP and a platelet count of $20 \times 10^9/L$ or less
457 received a total of 2 g/kg dose of Privigen administered as 1 g/kg infusions
458 daily for 2 consecutive days (*see Clinical Studies [14.2]*). Subjects ranged
459 in age from 15 to 69; 23 (40.4%) were male and 34 (59.6%) were female.

460

461 Concomitant medications affecting platelets or other treatments for
462 chronic ITP were not allowed. Thirty-two (56.1%) subjects received
463 premedication with acetaminophen and/or an antihistamine.

464

465
466 Table 6 lists the temporally associated AEs that occurred in >5% of
467 subjects with chronic ITP during a Privigen infusion or within 72 hours
468 after the end of a treatment cycle (two consecutive infusions) with
469 Privigen, *irrespective of causality*.

470
471 **Table 6: Chronic ITP Study – Adverse Events Occurring in >5% of**
472 **Subjects During a Privigen Infusion or Within 72 hours**
473 **After the End of a Treatment Cycle*, *Irrespective of***
474 ***Causality***

Adverse Event	Number (%) of Subjects [n=57]	Number (Rate) of Infusions With Adverse Event [n=114]
Headache	37 (64.9)	41 (0.360)
Pyrexia/hyperthermia	21 (36.8)	22 (0.193)
Nausea	6 (10.5)	6 (0.053)
Epistaxis	6 (10.5)	6 (0.053)
Vomiting	6 (10.5)	6 (0.053)
Blood unconjugated bilirubin increased	6 (10.5)	6 (0.053)
Blood conjugated bilirubin increased	5 (8.8)	5 (0.044)
Blood total bilirubin increased	4 (7.0)	4 (0.035)
Hematocrit decreased	3 (5.3)	3 (0.026)

476 * Two consecutive daily infusions.

477
478 Table 7 lists the adverse reactions (AEs considered to be “at least possibly
479 related” to the infusion of Privigen) that occurred in >5% of subjects,
480 *irrespective of time of occurrence.*

481
482 **Table 7: Chronic ITP Study – Adverse Reactions Considered “at Least
483 Possibly Related” to Privigen Occurring in >5% of Subjects,
484 Irrespective of Time of Occurrence**
485

Adverse Reaction	Number (%) of Subjects [n=57]	Number (Rate) of Infusions With Adverse Reaction [n=114]
Headache	37 (64.9)	52 (0.456)
Pyrexia/hyperthermia	19 (33.3)	21 (0.184)
Positive DAT	6 (10.5)	7 (0.061)
Anemia	6 (10.5)	6 (0.053)
Vomiting	5 (8.8)	6 (0.053)
Nausea	5 (8.8)	7 (0.061)
Bilirubin conjugated, increased	5 (8.8)	5 (0.044)
Bilirubin unconjugated, increased	5 (8.8)	5 (0.044)
Hyperbilirubinemia	3 (5.3)	3 (0.026)
Blood lactate dehydrogenase increased	3 (5.3)	3 (0.026)
Hematocrit decreased	3 (5.3)	3 (0.026)

486
487 Of the 149 non-serious AEs related to Privigen, 103 were mild, 37 were
488 moderate, and 9 were severe.

489
490 Three subjects experienced three SAEs, one of which (aseptic meningitis)
491 was related to the infusion of Privigen.

492
493 One subject withdrew from the study due to gingival bleeding that was not
494 related to Privigen.

495
496 Eight subjects, all of whom had a positive DAT, experienced transient
497 drug-related hemolytic reactions, which were associated with elevated
498 bilirubin, elevated lactate dehydrogenase, and a decrease in hemoglobin
499 level within two days after the infusion of Privigen. Two of the eight
500 subjects were clinically anemic but did not require clinical intervention;
501 these cases resolved uneventfully.

502
503 Four other subjects with active bleeding were reported to have developed
504 anemia without evidence of hemolysis.

505
506 In this study, there was a decrease in hemoglobin after the first Privigen
507 infusion (median decrease of 1.2 g/dL by Day 8) followed by a return to
508 near baseline by Day 29.

509

510 Fifty-six of the 57 subjects in this study had a negative DAT at baseline.
511 Of these 56 subjects, 12 (21.4%) developed a positive DAT during the 29-
512 day study period.

513

514 **6.2 Postmarketing Experience**

515 *Because adverse reactions are reported voluntarily post-approval from a*
516 *population of uncertain size, it is not always possible to reliably estimate the*
517 *frequency of these reactions or establish a causal relationship to product*
518 *exposure.*

519

520 **Privigen**

521 The following adverse reactions have been identified during
522 postmarketing use of Privigen:

- 523 • *Infusion reactions:* Hypersensitivity (e.g., anaphylaxis), changes in
524 blood pressure, dyspnea, chills and fever, tachycardia, chest
525 discomfort/pain, flushing
- 526 • *Hematologic:* Hemolytic anemia, jaundice/hyperbilirubinemia,
527 hemoglobinuria/hematuria/chromaturia, renal failure
- 528 • *Neurological:* Headache, aseptic meningitis, photophobia,
529 dizziness
- 530 • *Integumentary:* Urticaria, pruritus, rash

531

532 **General**

533 The following adverse reactions have been identified and reported during
534 the post-approval use of immune globulin products.¹⁴

535

- 536 • *Infusion Reactions:* Hypersensitivity (e.g., anaphylaxis), headache,
537 diarrhea, tachycardia, fever, fatigue, dizziness, malaise, chills,
538 flushing, urticaria or other skin reactions, wheezing or other chest
539 discomfort, nausea, vomiting, rigors, back pain, myalgia,
540 arthralgia, and changes in blood pressure
- 541 • *Renal:* Acute renal dysfunction/failure, osmotic nephropathy
- 542 • *Respiratory:* Apnea, Acute Respiratory Distress Syndrome
543 (ARDS), TRALI, cyanosis, hypoxemia, pulmonary edema,
544 dyspnea, bronchospasm
- 545 • *Cardiovascular:* Cardiac arrest, thromboembolism, vascular
546 collapse, hypotension
- 547 • *Neurological:* Coma, loss of consciousness, seizures, tremor,
548 aseptic meningitis syndrome
- 549 • *Integumentary:* Stevens-Johnson syndrome, epidermolysis,
550 erythema multiforme, bullous dermatitis
- 551 • *Hematologic:* Pancytopenia, leukopenia, hemolysis, positive DAT
552 (Coombs' test)
- 553 • *Musculoskeletal:* Back pain
- 554 • *Gastrointestinal:* Hepatic dysfunction, abdominal pain

- 555 • *General/Body as a Whole*: Pyrexia, rigors

556

557

558 **7 DRUG INTERACTIONS**

559

560 **7.1 Live Virus Vaccines**

561 The passive transfer of antibodies with immunoglobulin administration
562 may interfere with the response to live virus vaccines such as measles,
563 mumps, rubella, and varicella (*see Patient Counseling Information [17]*).¹⁵

564

565 Inform the immunizing physician of recent therapy with Privigen so that
566 appropriate measures can be taken.

567

568 **7.2 Serological Testing**

569 Various passively transferred antibodies in immunoglobulin preparation
570 may lead to misinterpretation of the results of serological testing.

571

572

573 **8 USE IN SPECIFIC POPULATIONS**

574

575 **8.1 Pregnancy**

576 Pregnancy Category C. Animal reproduction studies have not been
577 conducted with Privigen. It is not known whether Privigen can cause fetal
578 harm when administered to a pregnant woman or can affect reproduction
579 capacity. Privigen should be given to pregnant women only if clearly
580 needed. Immunoglobulins cross the placenta from maternal circulation
581 increasingly after 30 weeks of gestation.^{16,17}

582

583 **8.3 Nursing Mothers**

584 Use of Privigen in nursing mothers has not been evaluated.

585

586 **8.4 Pediatric Use**

587 Treatment of Primary Humoral Immunodeficiency

588 Privigen was evaluated in 31 pediatric subjects (19 children and 12
589 adolescents) with PI (pivotal study). There were no apparent differences
590 in the safety and efficacy profiles as compared to those in adult subjects.
591 No pediatric-specific dose requirements were necessary to achieve the
592 desired serum IgG levels. The safety and effectiveness of Privigen have
593 not been established in pediatric patients with PI who are under the age
594 of 3.

595

596 Treatment of Chronic Immune Thrombocytopenic Purpura

597 The safety and effectiveness of Privigen have not been established in
598 pediatric patients with chronic ITP who are under the age of 15.

599

600 8.5 Geriatric Use

601 Clinical studies of Privigen did not include sufficient numbers of subjects
602 age 65 and over to determine whether they respond differently from
603 younger subjects.

604
605 Use caution when administering Privigen to patients age 65 and over who
606 are judged to be at increased risk of developing acute renal insufficiency
607 and thrombotic events (*see Boxed Warning, Warnings and Precautions*
608 *[5.2, 5.3]*). Do not exceed recommended doses, and administer Privigen
609 at the minimum infusion rate practicable.

610
611

612 10 OVERDOSAGE

613
614 Overdose may lead to fluid overload and hyperviscosity, particularly in
615 the elderly and in patients with impaired renal function.

616
617

618 11 DESCRIPTION

619

620 Privigen is a ready-to-use, sterile, 10% protein liquid preparation of
621 polyvalent human immunoglobulin G (IgG) for intravenous
622 administration. Privigen has a purity of at least 98% IgG, consisting
623 primarily of monomers. The balance consists of IgG dimers ($\leq 12\%$),
624 small amounts of fragments and polymers, and albumin. Privigen contains
625 ≤ 25 mcg/mL IgA. The IgG subclass distribution (approximate mean
626 values) is IgG₁, 67.8%; IgG₂, 28.7%; IgG₃, 2.3%; and IgG₄, 1.2%.
627 Privigen has an osmolality of approximately 320 mOsmol/kg (range: 240
628 to 440) and a pH of 4.8 (range: 4.6 to 5.0).

629

630 Privigen contains approximately 250 mmol/L (range: 210 to 290) of L-
631 proline (a nonessential amino acid) as a stabilizer and trace amounts of
632 sodium. Privigen contains no carbohydrate stabilizers (e.g., sucrose,
633 maltose) and no preservative.

634

635 Privigen is prepared from large pools of human plasma by a combination
636 of cold ethanol fractionation, octanoic acid fractionation, and anion
637 exchange chromatography. The IgG proteins are not subjected to heating
638 or to chemical or enzymatic modification. The Fc and Fab functions of
639 the IgG molecule are retained. Fab functions tested include antigen
640 binding capacities, and Fc functions tested include complement activation
641 and Fc-receptor-mediated leukocyte activation (determined with
642 complexed IgG). Privigen does not activate the complement system or
643 prekallikrein in an unspecific manner.

644

645 All plasma units used in the manufacture of Privigen have been tested and
646 approved for manufacture using FDA-licensed serological assays for
647 hepatitis B surface antigen and antibodies to HCV and HIV-1/2 as well as
648 FDA-licensed Nucleic Acid Testing (NAT) for HCV and HIV-1 and found
649 to be nonreactive (negative). For HBV, an investigational NAT procedure
650 is used and the plasma units found to be negative; however, the
651 significance of a negative result has not been established. In addition, the
652 plasma has been tested for B19 virus (B19V) DNA by NAT. Only plasma
653 that passed virus screening is used for production, and the limit for B19V
654 in the fractionation pool is set not to exceed 10^4 IU of B19V DNA per mL.
655

656 The manufacturing process for Privigen includes three steps to reduce the
657 risk of virus transmission. Two of these are dedicated virus clearance
658 steps: pH 4 incubation to inactivate enveloped viruses and virus filtration
659 to remove, by size exclusion, both enveloped and non-enveloped viruses
660 as small as approximately 20 nanometers. In addition, a depth filtration
661 step contributes to the virus reduction capacity.
662

663 These steps have been independently validated in a series of *in vitro*
664 experiments for their capacity to inactivate and/or remove both enveloped
665 and non-enveloped viruses.
666

667 Table 8 shows the virus clearance during the manufacturing process for
668 Privigen, expressed as the mean \log_{10} reduction factor (LRF).
669

670 **Table 8: Virus Inactivation/Removal in Privigen***
671

	HIV-1	PRV	BVDV	WNV	EMCV	MVM
Virus property						
Genome	RNA	DNA	RNA	RNA	RNA	DNA
Envelope	Yes	Yes	Yes	Yes	No	No
Size (nm)	80-100	120-200	50-70	50-70	25-30	18-24
Manufacturing step	Mean LRF					
pH 4 incubation	≥ 5.4	≥ 5.9	4.6	≥ 7.8	nt	nt
Depth filtration	≥ 5.3	≥ 6.3	2.1	3.0	4.2	2.3
Virus filtration	≥ 5.3	≥ 5.5	≥ 5.1	≥ 5.9	≥ 5.4	≥ 5.5
Overall reduction (\log_{10} units)	≥ 16.0	≥ 17.7	≥ 11.8	≥ 16.7	≥ 9.6	≥ 7.8

672 HIV-1, human immunodeficiency virus type 1, a model for HIV-1 and HIV-2; PRV, pseudorabies
673 virus, a nonspecific model for large enveloped DNA viruses (e.g., herpes virus); BVDV, bovine
674 viral diarrhea virus, a model for hepatitis C virus; WNV, West Nile virus; EMCV,
675 encephalomyocarditis virus, a model for hepatitis A virus; MVM, minute virus of mice, a model for
676 a small highly resistant non-enveloped DNA virus (e.g., parvovirus); LRF, \log_{10} reduction factor;
677 nt, not tested.

678 * The virus clearance of human parvovirus B19 was investigated experimentally at the pH 4
679 incubation step. The estimated LRF obtained was ≥ 5.3 .

680

681 The manufacturing process was also investigated for its capacity to
682 decrease the infectivity of an experimental agent of transmissible
683 spongiform encephalopathy (TSE), considered a model for CJD and its
684 variant vCJD.¹⁸ Several of the production steps have been shown to
685 decrease TSE infectivity of an experimental model agent. TSE reduction
686 steps include octanoic acid fractionation ($\geq 6.4 \log_{10}$), depth filtration (2.6
687 \log_{10}), and virus filtration ($\geq 5.8 \log_{10}$). These studies provide reasonable
688 assurance that low levels of vCJD/CJD agent infectivity, if present in the
689 starting material, would be removed.

690

691

692 12 CLINICAL PHARMACOLOGY

693

694 12.1 Mechanism of Action

695 Treatment of Primary Humoral Immunodeficiency

696 Privigen is a replacement therapy for primary humoral immunodeficiency,
697 and supplies a broad spectrum of opsonic and neutralizing IgG antibodies
698 against bacterial, viral, parasitic and mycoplasma agents and their toxins.
699 The mechanism of action in PI has not been fully elucidated.

700

701 Treatment of Chronic Immune Thrombocytopenic Purpura

702 The mechanism of action of high doses of immunoglobulins in the
703 treatment of chronic ITP has not been fully elucidated.

704

705 12.3 Pharmacokinetics

706 Treatment of Primary Humoral Immunodeficiency

707 In the clinical study (pivotal study) assessing the efficacy and safety of
708 Privigen in 80 subjects with PI (*see Clinical Studies [14.1]*), serum
709 concentrations of total IgG and IgG subclasses were measured in 25
710 subjects (ages 13 to 69) following the 7th infusion for the 3 subjects on the
711 3-week dosing interval and following the 5th infusion for the 22 subjects
712 on the 4-week dosing interval. The dose of Privigen used in these subjects
713 ranged from 200.0 mg/kg to 714.3 mg/kg. After the infusion, blood
714 samples were taken until Day 21 and Day 28 for the 3-week and 4-week
715 dosing intervals, respectively.

716

717
718 Table 9 summarizes the pharmacokinetic parameters of Privigen, based on
719 serum concentrations of total IgG.

720
721 **Table 9: PI Pivotal Study – Pharmacokinetic Parameters of Privigen**
722 **in Subjects**
723

Parameter	3-Week Dosing Interval (n=3)		4-Week Dosing Interval (n=22)	
	Mean (SD)	Median (Range)	Mean (SD)	Median (Range)
C _{max} (peak, mg/dL)	2,550 (400)	2,340 (2,290-3,010)	2,260 (530)	2,340 (1,040-3,460)
C _{min} (trough, mg/dL)	1,230 (230)	1,200 (1,020-1,470)	1,000 (200)	1,000 (580-1,360)
t _{1/2} (days)	27.6 (5.9)	27.8 (21.6-33.4)	45.4 (18.5)	37.3 (20.6-96.6)
AUC _{0-t} (day × mg/dL)*	32,820 (6,260)	29,860 (28,580-40,010)	36,390 (5,950)	36,670 (19,680-44,340)
AUC _{0-∞} (day × mg/dL) *	79,315 (20,170)	78,748 (59,435-99,762)	104,627 (33,581)	98,521 (64,803-178,600)
Clearance (mL/day/kg)*	1.3 (0.1)	1.3 (1.1-1.4)	1.3 (0.3)	1.3 (0.9-2.1)
Mean residence time (days) *	38.6 (8.1)	39.5 (30.1-46.2)	65.2 (24.7)	59.0 (33.2-129.6)
Volume of distribution at steady state (mL/kg) *	50 (13)	44 (40-65)	84 (35)	87 (40-207)

724 C_{max}, maximum serum concentration; C_{min}, trough (minimum level) serum concentration;
725 t_{1/2}, elimination half-life; AUC_{0-t}, area under the curve from 0 hour to last sampling time;
726 AUC_{0-∞}, area under the curve from 0 hour to infinite time.

727 * Calculated by log-linear trapezoidal rule.

728
729 The median half-life of Privigen was 36.6 days for the 25 subjects in the
730 pharmacokinetic subgroup.

731
732 Although no systematic study was conducted to evaluate the effect of
733 gender and age on the pharmacokinetics of Privigen, based on the small
734 sample size (11 males and 14 females) it appears that clearance of
735 Privigen is comparable in males (1.27 ± 0.35 mL/day/kg) and females
736 (1.34 ± 0.22 mL/day/kg). In six subjects between 13 and 15 years of age,
737 the clearance of Privigen (1.35 ± 0.44 mL/day/kg) is comparable to that
738 observed in 19 adult subjects 19 years of age or older (1.29 ± 0.22
739 mL/day/kg).

740

741 The IgG subclass levels observed in the pharmacokinetic study were
742 consistent with a physiologic distribution pattern (mean trough values):
743 IgG₁, 564.91 mg/dL; IgG₂, 394.15 mg/dL; IgG₃, 30.16 mg/dL; IgG₄,
744 10.88 mg/dL.

745

746 Treatment of Chronic Immune Thrombocytopenic Purpura

747 Pharmacokinetic studies with Privigen were not performed in subjects
748 with chronic ITP.

749

750

751 **14 CLINICAL STUDIES**

752

753 **14.1 Treatment of Primary Humoral Immunodeficiency**

754 A prospective, open-label, single-arm, multicenter study (pivotal study)
755 assessed the efficacy, safety, and pharmacokinetics of Privigen in adult
756 and pediatric subjects with PI, who were treated for 12 months at a 3-week
757 or 4-week dosing interval. Subjects ranged in age from 3 to 69; 46
758 (57.5%) were male and 34 (42.5%) were female; 77.5% were Caucasian,
759 15% were Hispanic, and 7.5% were African-American. All subjects had
760 been on regular IGIV replacement therapy for at least 6 months prior to
761 participating in the study.

762

763 The efficacy analysis included 80 subjects, 16 (20%) on the 3-week dosing
764 interval and 64 (80%) on the 4-week dosing interval. Doses ranged from
765 200 mg/kg to 888 mg/kg per infusion. The median dose for the 3-week
766 interval was 428.3 mg/kg per infusion; the median dose for the 4-week
767 interval was 440.6 mg/kg per infusion. Subjects received a total of 1038
768 infusions of Privigen, 272 for the 3-week dosing regimen and 766 for the
769 4-week dosing regimen. The maximum infusion rate allowed during this
770 study was 8 mg/kg/min with 715 (69%) of the infusions administered at a
771 rate of 7 mg/kg/min or greater.

772

773 The primary analysis for efficacy was based on the annual rate of acute
774 serious bacterial infections (aSBI), defined as pneumonia,
775 bacteremia/septicemia, osteomyelitis/septic arthritis, bacterial meningitis,
776 and visceral abscess, per subject per year. Secondary analyses were based
777 on the annual rate of other infections, antibiotic use, days out of
778 work/school/day care or unable to perform normal activities due to illness,
779 and days of hospitalization.

780

781 During the 12-month study period, the aSBI rate was 0.08 (with an upper
782 1-sided 99% confidence interval of 0.203), which met the predefined
783 success rate of less than one aSBI per subject per year. Six subjects
784 experienced an aSBI, including three cases of pneumonia and one case

785 each of septic arthritis, osteomyelitis, and visceral abscess. All six
786 subjects completed the study.

787
788 The rate of other infections was 3.55 infections per subject per year. The
789 infections that occurred most frequently were sinusitis (31.3%),
790 nasopharyngitis (22.5%), upper respiratory tract infection (18.8%),
791 bronchitis (13.8%), and rhinitis (13.8%). Among the 255 infections, 16
792 (6.3%) occurring in 10 subjects were considered severe.

793
794 Table 10 summarizes the efficacy results for all 80 subjects.

795
796 **Table 10: PI Pivotal Study – Summary of Efficacy Results in Subjects**
797

Number of Subjects	80
Results from Case Report Forms	
Total Number of Subject Days	26,198
Infections	
Annual rate of confirmed aSBI*s†	0.08 aSBIs/subject year [†]
Annual rate of other infections	3.55 infections/subject year
Antibiotic use	
Number of subjects (%)	64 (80%)
Annual rate	87.4 days/subject year
Results from Subject Diaries	
Total Number of Diary Days	24,059
Out of work/school/day care or unable to perform normal activities due to illness	
Number of days (%)	570 (2.37%)
Annual rate	8.65 days/subject year
Hospitalization	
Number of days (%)	166 (0.69%)
Annual rate	2.52 days/subject year

798 * Defined as pneumonia, bacterial meningitis, bacteremia/septicemia, osteomyelitis/septic arthritis,
799 and visceral abscess.

800 † Upper 1-sided 99% confidence interval: 0.203.

801
802 **14.2 Treatment of Chronic Immune Thrombocytopenic Purpura**

803 A prospective, open-label, single-arm, multicenter study assessed the
804 efficacy, safety, and tolerability of Privigen in 57 subjects with chronic
805 ITP and a platelet count of $20 \times 10^9/L$ or less. Subjects ranged in age from
806 15 to 69; 23 (40.4%) were male and 34 (59.6%) were female; all were
807 Caucasian.

808
809 Subjects received a 2 g/kg dosage of Privigen administered as 1 g/kg
810 (10 mL/kg) intravenous infusion daily for 2 consecutive days, and were
811 observed for 29 days. Fifty-three (93%) subjects received Privigen at the
812 maximum infusion rate allowed (4 mg/kg/min [0.04 mL/kg/min]).

813

814 The primary analysis was based on the response rate defined as the
815 percentage of subjects with an increase in platelet counts to at least $50 \times 10^9/L$
816 within 7 days after the first infusion (responders). Secondary
817 analyses were based on the increase in platelet counts and the time to
818 reach a platelet count of at least $50 \times 10^9/L$ at any point within the study
819 period, the duration of that response, and the regression (decrease in the
820 severity) of hemorrhage in subjects who had bleeding at baseline. Platelet
821 counts were measured on Days 1, 2, 4, 6, 8, 15, 22, and 29. Additional
822 measurements on Days 57 and 85 occurred in subjects with a platelet
823 count of at least $50 \times 10^9/L$ at the previous visit.

824
825 Of the 57 subjects in the efficacy analysis, 46 (80.7%) responded to
826 Privigen with a rise in platelet counts to at least $50 \times 10^9/L$ within 7 days
827 after the first infusion. The lower bound of the 95% confidence interval
828 for the response rate (69.2%) is above the predefined response rate of
829 50%.

830
831 The highest median increase in platelet counts was seen 7 days after the
832 first infusion ($123 \times 10^9/L$). The median maximum platelet count
833 achieved was $154 \times 10^9/L$. The median time to reach a platelet response
834 of more than $50 \times 10^9/L$ was 2.5 days after the first infusion. Twenty-five
835 (43%) of the 57 subjects reached this response by Day 2 prior to the
836 second infusion and 43 (75%) subjects reached this response by Day 6.

837
838 The duration of platelet response was analyzed for the 48 subjects who
839 achieved a response any time after the first infusion. The median duration
840 of platelet response in these subjects was 15.4 days (range: 1 to >82 days).
841 Thirty-six (75%) of the 48 subjects maintained the response for at least 8.8
842 days and 12 (25%) of them for at least 21.9 days. Five (9%) subjects
843 maintained a response up to Day 29 and two (4%) up to Day 85.

844
845 A decrease in the severity of hemorrhage from baseline was observed in
846 the following bleeding locations: skin (31 of 36 subjects), oral cavity (11
847 of 11 subjects), and genitourinary tract (7 of 9 subjects). This decrease
848 was not sustained in all subjects up to the end of the 29-day study period.

849

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16 HOW SUPPLIED/STORAGE AND HANDLING

908 Privigen is supplied in a single-use, tamper-evident vial containing the
909 labeled amount of functionally active IgG. The components used in the
910 packaging for Privigen are latex-free.

911
912 The following presentations of Privigen are available:
913

NDC Number	Fill Size (mL)	Grams Protein
44206-436-05	50	5
44206-437-10	100	10
44206-438-20	200	20

914
915
916
917

Each vial has an integral suspension band and a label with two peel-off strips showing the product name, lot number, and expiration date.

918 When stored at room temperature (up to 25°C [77°F]), Privigen is stable
919 for up to 36 months, as indicated by the expiration date printed on the
920 outer carton and vial label.

921
922 Keep Privigen in its original carton to protect it from light.

923
924 Do not freeze.

925
926

17 PATIENT COUNSELING INFORMATION

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Inform patients of the early signs of hypersensitivity reactions to Privigen (including hives, generalized urticaria, tightness of the chest, wheezing, hypotension, and anaphylaxis), and advise them to notify their physician if they experience any of these symptoms.

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Inform patients to immediately report the following signs and symptoms to their physician:

- 936 • Decreased urine output, sudden weight gain, fluid retention/edema,
937 and/or shortness of breath, which may suggest kidney problems
- 938 • Shortness of breath, changes in mental status, chest pain, and other
939 manifestations of thrombotic events
- 940 • Severe headache, neck stiffness, drowsiness, fever, sensitivity to
941 light, painful eye movements, nausea, and vomiting, which may
942 suggest aseptic meningitis syndrome
- 943 • Fatigue, increased heart rate, yellowing of skin or eyes, and dark-
944 colored urine, which may suggest hemolysis

- 945 • Severe breathing problems, lightheadedness, drops in blood pressure,
946 and fever, which may suggest TRALI (a condition typically
947 occurring within 1 to 6 hours following transfusion)
948

949 Inform patients that Privigen is made from human blood and may contain
950 infectious agents that can cause disease (e.g., viruses and, theoretically the
951 CJD agent). Explain that the risk that Privigen may transmit an infectious
952 agent has been reduced by screening the plasma donors, by testing donated
953 plasma for certain virus infections, and by inactivating or removing certain
954 viruses during manufacturing, and counsel patients to report any
955 symptoms that concern them.
956

957 Inform patients that administration of IgG may interfere with the response
958 to live virus vaccines (e.g., measles, mumps, rubella, and varicella), and
959 instruct them to notify their immunizing physician of recent therapy with
960 Privigen.
961

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