

1. NAME OF THE MEDICINAL PRODUCT

Taltz 80 mg solution for injection in pre-filled syringe.

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each pre-filled syringe contains 80 mg ixekizumab in 1 ml.

Ixekizumab is produced in CHO cells by recombinant DNA technology.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Solution for injection (injection).

The solution is clear and colourless to slightly yellow.

4. CLINICAL PARTICULARS**4.1 Therapeutic indications****Plaque psoriasis**

Taltz is indicated for the treatment of moderate to severe plaque psoriasis in adults who are candidates for systemic therapy.

Paediatric plaque psoriasis

Taltz is indicated for the treatment of moderate to severe plaque psoriasis in children from the age of 6 years and adolescents who are candidates for systemic therapy.

Psoriatic arthritis

Taltz, alone or in combination with methotrexate, is indicated for the treatment of active psoriatic arthritis in adult patients who have responded inadequately to, or who are intolerant to one or more disease-modifying anti-rheumatic drug (DMARD) therapies (see section 5.1).

Axial spondyloarthritis***Ankylosing spondylitis (radiographic axial spondyloarthritis)***

Taltz is indicated for the treatment of adult patients with active ankylosing spondylitis who have responded inadequately to conventional therapy.

Non-radiographic axial spondyloarthritis

Taltz is indicated for the treatment of adult patients with active non-radiographic axial spondyloarthritis with objective signs of inflammation as indicated by elevated C-reactive protein (CRP) and/or magnetic resonance imaging (MRI) who have responded inadequately to nonsteroidal anti-inflammatory drugs (NSAIDs).

4.2 Posology and method of administration

This medicinal product is intended for use under the guidance and supervision of a physician experienced in the diagnosis and treatment of conditions for which it is indicated.

58 Posology

59 *Plaque psoriasis in adults*

60 The recommended dose is 160 mg by subcutaneous injection (two 80 mg injections) at week 0,
61 followed by 80 mg (one injection) at weeks 2, 4, 6, 8, 10, and 12, then maintenance dosing of 80 mg
62 (one injection) every 4 weeks (Q4W).

64 *Paediatric plaque psoriasis (age 6 years and above)*

65 Efficacy and safety data is not available in children below the age of 6 years (see section 5.1).

66
67 TALTZ is administered by subcutaneous injection every 4 weeks (Q4W). The recommended dose in
68 pediatric patients from 6 to less than 18 years of age with moderate-to-severe plaque psoriasis is based
69 on the following weight categories.

Children's body weight	Recommended starting dose (week 0)	Recommended dose every 4 weeks (Q4W) thereafter
Greater than 50 kg	160 mg (two 80 mg injections)	80 mg
25 to 50 kg	80 mg	40 mg
Less than 25 kg	40 mg	20 mg

71
72 For children prescribed 80 mg, Taltz can be used directly from the prefilled syringe.
73 For instructions on preparation of Taltz 40 mg, see section 6.6. Doses less than 80 mg must be
74 prepared by a healthcare professional.
75 Taltz is not recommended for use in children with a body weight below 25 kg. Paediatric body weights
76 must be recorded and regularly re-checked prior to dosing.

78 *Psoriatic arthritis*

79 The recommended dose is 160 mg by subcutaneous injection (two 80 mg injections) at week 0,
80 followed by 80 mg (one injection) every 4 weeks thereafter. For psoriatic arthritis patients with
81 concomitant moderate to severe plaque psoriasis, the recommended dosing regimen is the same as for
82 plaque psoriasis.

84 *Axial spondyloarthritis (radiographic and non-radiographic)*

85 The recommended dose is 160 mg (two 80 mg injections) by subcutaneous injection at week 0,
86 followed by 80 mg every 4 weeks (see section 5.1 for further information).

88 For all indications (plaque psoriasis in adults and children, psoriatic arthritis, axial spondyloarthritis)
89 consideration should be given to discontinuing treatment in patients who have shown no response after
90 16 to 20 weeks of treatment. Some patients with initially partial response may subsequently improve
91 with continued treatment beyond 20 weeks.

93 Special populations

95 *Elderly (≥ 65 years)*

96 No dose adjustment is required (see section 5.2).

98 There is limited information in subjects aged ≥ 75 years.

100 *Renal or hepatic impairment*

101 Taltz has not been studied in these patient populations. No dose recommendations can be made.

103 Paediatric population

105 *Paediatric plaque psoriasis (below a body weight of 25 kg and below the age of 6 years)*

106 There is no relevant use of Taltz in children below a body weight of 25 kg and below the age of
107 6 years in the treatment of moderate to severe plaque psoriasis.

109 *Paediatric psoriatic arthritis*

110 The safety and efficacy of Taltz in children and adolescents aged 2 to less than 18 years in the
111 treatment of psoriatic arthritis (a category of juvenile idiopathic arthritis) have not yet been
112 established. No data are available.
113 There is no relevant use of Taltz in children below 2 years for the indication of psoriatic arthritis.
114

115 Method of administration

116
117 Subcutaneous use.

118 Taltz is for subcutaneous injection. Injection sites may be alternated. If possible, areas of the skin that
119 show psoriasis should be avoided as injection sites. The solution/the syringe must not be shaken.
120

121 After proper training in subcutaneous injection technique, patients may self-inject Taltz if a healthcare
122 professional determines that it is appropriate. However, the physician should ensure appropriate
123 follow-up of patients. Comprehensive instructions for administration are given in the package leaflet
124 and the user manual.

125
126 Doses less than 80 mg which require dose preparation should only be administered by a healthcare
127 professional.
128

129 For instructions on preparation of the medicinal product before administration, see section 6.6.
130

131 **4.3 Contraindications**

132
133 Serious hypersensitivity to the active substance or to any of the excipients listed in section 6.1.
134

135 Clinically important active infections (e.g. active tuberculosis, see section 4.4).
136

137 **4.4 Special warnings and precautions for use**

138 139 Traceability

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141 In order to improve the traceability of biological medicinal products, the name and the batch number
142 of the administered product should be clearly recorded.
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145 Infections

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147 Treatment with Taltz is associated with an increased rate of infections such as upper respiratory tract
148 infection, oral candidiasis, conjunctivitis, and tinea infections (see section 4.8).
149

150 Taltz should be used with caution in patients with clinically important chronic infection or history of
151 recurrent infection. Patients should be instructed to seek medical advice if sign or symptoms
152 suggestive of an infection occur. If an infection develops, patients should be carefully monitored and
153 Taltz discontinued if the patient is not responding to standard therapy or the infection becomes serious.
154 Taltz should not be resumed until the infection resolves.
155

156 Taltz must not be given to patients with active tuberculosis (TB). Anti-TB therapy prior to initiation of
157 Taltz in patients with latent TB should be considered.
158

159 Hypersensitivity

160
161 Serious hypersensitivity reactions, including some cases of anaphylaxis, angioedema, urticaria and,
162 rarely, late (10-14 days following injection) serious hypersensitivity reactions including widespread
163 urticaria, dyspnea and high antibody titres have been reported. If a serious hypersensitivity reaction
164 occurs, administration of Taltz should be discontinued immediately and appropriate therapy initiated.
165

166 Inflammatory bowel disease (including Crohn's disease and ulcerative colitis)

167
168 Cases of new or exacerbations of inflammatory bowel disease have been reported with ixekizumab
169 (see section 4.8). Ixekizumab is not recommended in patients with inflammatory bowel disease. If a
170 patient develops signs and symptoms of inflammatory bowel disease or experiences an exacerbation of
171 pre-existing inflammatory bowel disease, ixekizumab should be discontinued and appropriate medical
172 management should be initiated.

173
174 Immunisations

175
176 Taltz should not be used with live vaccines. No data are available on the response to live vaccines;
177 there are insufficient data on response to inactive vaccines (see section 5.1).

178
179 Excipients

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181 This medicinal product contains less than 1 mmol sodium (23 mg) per 80 mg dose, that is to
182 say essentially "sodium-free".

183
184 **4.5 Interaction with other medicinal products and other forms of interaction**

185
186 In plaque psoriasis studies, the safety of Taltz in combination with other immunomodulatory agents or
187 phototherapy has not been evaluated.

188 In population pharmacokinetic analyses, clearance of ixekizumab was not affected by concomitant
189 administration of oral corticosteroids, NSAIDs, sulfasalazine, or methotrexate.

190
191 Cytochrome P450 substrates

192
193 Results from an interaction study in patients with moderate-to-severe psoriasis determined that
194 12 weeks of administration of ixekizumab with substances metabolised by CYP3A4 (i.e., midazolam),
195 CYP2C9 (i.e., warfarin), CYP2C19 (i.e., omeprazole), CYP1A2 (i.e., caffeine) or CYP2D6 (i.e.,
196 dextromethorphan) does not have a clinically significant impact on the pharmacokinetics of these
197 substances.

198
199 **4.6 Fertility, pregnancy and lactation**

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201 Women of childbearing potential

202
203 Women of childbearing potential should use an effective method of contraception during treatment
204 and for at least 10 weeks after treatment.

205
206 Pregnancy

207
208 There is a limited amount of data from the use of ixekizumab in pregnant women. Animal studies do
209 not indicate direct or indirect harmful effects with respect to pregnancy, embryonic/foetal
210 development, parturition or post-natal development (see section 5.3). As a precautionary measure, it is
211 preferable to avoid the use of Taltz during pregnancy.

212
213 Breast-feeding

214
215 It is not known whether ixekizumab is excreted in human milk or absorbed systemically after
216 ingestion. However, ixekizumab is excreted at low levels in the milk of cynomolgus monkeys. A
217 decision should be made whether to discontinue breast-feeding or to discontinue Taltz taking into
218 account the benefit of breast-feeding for the child and the benefit of therapy for the woman.

219

220 Fertility

221
222 The effect of ixekizumab on human fertility has not been evaluated. Animal studies do not indicate
223 direct or indirect harmful effects with respect to fertility (see section 5.3).
224

225 **4.7 Effects on ability to drive and use machines**

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227 Taltz has no or negligible influence on the ability to drive and use machines.
228

229 **4.8 Undesirable effects**

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231 Summary of the safety profile

232
233 The most frequently reported adverse reactions were injection site reactions (15.5%) and upper
234 respiratory tract infections (16.4%) (most frequently nasopharyngitis).
235

236 Tabulated list of adverse reactions

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238 Adverse reactions from clinical studies and postmarketing reports (Table 1) are listed by MedDRA
239 system organ class. Within each system organ class, the adverse reactions are ranked by frequency,
240 with the most frequent reactions first. Within each frequency grouping, adverse reactions are presented
241 in order of decreasing seriousness. In addition, the corresponding frequency category for each adverse
242 reactions is based on the following convention: very common ($\geq 1/10$); common ($\geq 1/100$ to $< 1/10$);
243 uncommon ($\geq 1/1,000$ to $< 1/100$); rare ($\geq 1/10,000$ to $< 1/1,000$); very rare ($< 1/10,000$).
244

245 A total of 8,956 patients have been treated with Taltz in blinded and open-label clinical studies in
246 plaque psoriasis, psoriatic arthritis, axial spondyloarthritis and other autoimmune conditions. Of these,
247 6,385 patients were exposed to Taltz for at least one year, cumulatively representing 19,833 adult
248 patient years of exposure and 196 children cumulatively representing 207 patient years of exposure.
249

250
251 **Table 1. List of adverse reactions in clinical studies^a and postmarketing reports**
252

System organ class	Frequency	Adverse reaction
Infections and infestations	Very common	Upper respiratory tract infection
	Common	Tinea infection, Herpes simplex (mucocutaneous)
	Uncommon	Influenza, Rhinitis, Oral candidiasis, Conjunctivitis, Cellulitis
Blood and lymphatic system disorders	Uncommon	Neutropenia, Thrombocytopenia
Immune system disorders	Uncommon	Angioedema
	Rare	Anaphylaxis
Respiratory, thoracic and mediastinal disorders	Common	Oropharyngeal pain
Gastrointestinal disorders	Common	Nausea
	Uncommon	Inflammatory bowel disease
Skin and subcutaneous disorders	Uncommon	Urticaria, Rash, Eczema
General disorders and administration site conditions	Very common	Injection site reactions ^a

253 ^a See section description of selected adverse reactions

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Description of selected adverse reactions

Injection site reactions

The most frequent injection site reactions observed were erythema and pain. These reactions were predominantly mild to moderate in severity and did not lead to discontinuation of Taltz.

In the adult plaque psoriasis studies, injection site reactions were more common in subjects with a body weight < 60 kg compared with the group with a body weight \geq 60 kg (25 % vs. 14 % for the combined Q2W and Q4W groups). In the psoriatic arthritis studies, injection site reactions were more common in subjects with a body weight < 100 kg compared with the group with a body weight \geq 100 kg (24 % vs. 13 % for the combined Q2W and Q4W groups). In the axial spondyloarthritis studies, injection site reactions were similar in subjects with a body weight < 100 kg compared with the group with a body weight \geq 100 kg (14 % vs. 9 % for the combined Q2W and Q4W groups). The increased frequency of injection site reactions in the combined Q2W and Q4W groups did not result in an increase in discontinuations in either the plaque psoriasis, the psoriatic arthritis or the axial spondyloarthritis studies.

Infections

In the placebo-controlled period of the phase III clinical studies in plaque psoriasis in adults, infections were reported in 27.2 % of patients treated with Taltz for up to 12 weeks compared with 22.9 % of patients treated with placebo.

The majority of infections were non-serious and mild to moderate in severity, most of which did not necessitate treatment discontinuation. Serious infections occurred in 13 (0.6 %) of patients treated with Taltz and in 3 (0.4 %) of patients treated with placebo (see section 4.4). Over the entire treatment period infections were reported in 52.8 % of patients treated with Taltz (46.9 per 100 patient years). Serious infections were reported in 1.6 % of patients treated with Taltz (1.5 per 100 patient years).

Infection rates observed in psoriatic arthritis and axial spondyloarthritis clinical studies were similar to those observed in the plaque psoriasis studies with the exception of the frequencies of the adverse reactions of influenza and conjunctivitis which were common in patients with psoriatic arthritis.

Laboratory assessment of neutropenia and thrombocytopenia

In plaque psoriasis studies, 9% of patients receiving Taltz developed neutropenia. In most cases, the blood neutrophil count was \geq 1,000 cells/mm³. Such levels of neutropenia may persist, fluctuate or be transient. 0.1% of patients receiving Taltz developed a neutrophil count <1000 cells/mm³. In general, neutropenia did not require discontinuation of Taltz. 3% of patients exposed to Taltz had a shift from a normal baseline platelet value to <150,000 platelet cells/mm³ to \geq 75,000 cells/mm³.

Thrombocytopenia may persist, fluctuate or be transient.

The frequency of neutropenia and thrombocytopenia in psoriatic arthritis and axial spondyloarthritis clinical studies is similar to that observed in the plaque psoriasis studies.

Immunogenicity

Approximately 9–17% of adult plaque psoriasis patients treated with Taltz at the recommended dosing regimen developed anti-drug antibodies, the majority of which were low titres and not associated with reduced clinical response up to 60 weeks of treatment. However, approximately 1% of patients treated with Taltz had confirmed neutralising antibodies associated with low drug concentrations and reduced clinical response.

In psoriatic arthritis patients treated with Taltz at the recommended dosing regimen up to 52 weeks, approximately 11% developed anti-drug antibodies, the majority of which were low titre, and

307 approximately 8% had confirmed neutralising antibodies. No apparent association between the
308 presence of neutralising antibodies and impact on drug concentration or efficacy was observed.
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310 In paediatric psoriasis patients treated with Taltz at the recommended dosing regimen up to 12 weeks,
311 21 patients (18%) developed anti-drug antibodies, approximately half were low titer and 5 patients
312 (4%) had confirmed neutralizing antibodies associated with low drug concentrations. There was no
313 association with clinical response or adverse events.
314

315 In radiographic axial spondyloarthritis patients treated with Taltz at the recommended dosing regimen
316 up to 16 weeks, 5.2% developed anti-drug antibodies, the majority of which were low titer, and 1.5%
317 (3 patients) had neutralising antibodies (NAb). In these 3 patients, NAb-positive samples had low
318 ixekizumab concentrations and none of these patients achieved an ASAS40 response. In
319 non-radiographic axial spondyloarthritis patients treated with Taltz at the recommended dosing
320 regimen for up to 52 weeks, 8.9% developed anti-drug antibodies, all of which were low titer; no
321 patient had neutralising antibodies; and no apparent association between the presence of anti-drug
322 antibodies and drug concentration, efficacy, or safety was observed.
323

324 Across all indications, an association between immunogenicity and treatment emergent adverse events
325 has not been clearly established.
326

327 Paediatric population

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329 The safety profile observed in children with plaque psoriasis treated with Taltz every 4 weeks is
330 consistent with the safety profile in adult patients with plaque psoriasis with the exception of the
331 frequencies of conjunctivitis, influenza, and urticaria which were common. Inflammatory bowel
332 disease was also more frequent in paediatric patients, although it was still uncommon. In the paediatric
333 clinical study, Crohn's disease occurred in 0.9% of patients in the Taltz group and 0% of patients in
334 the placebo group during the 12-week, placebo-controlled period. Crohn's disease occurred in a total
335 of 4 Taltz treated subjects (2.0%) during the combined placebo-controlled and maintenance periods of
336 the paediatric clinical study.
337

338 Reporting of suspected adverse reactions

339 Reporting suspected adverse reactions after authorisation of the medicinal product is important. It
340 allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare
341 professionals are asked to report any suspected adverse reactions via the national reporting system
342 listed in Appendix V.
343

344 **4.9 Overdose**

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346 Doses up to 180 mg have been administered subcutaneously in clinical trials without dose-limiting
347 toxicity. Overdoses up to 240 mg, subcutaneously, as a single administration in clinical trials, have
348 been reported without any serious adverse events.

349 In the event of overdose, it is recommended that the patient be monitored for any signs or symptoms
350 of adverse reactions and appropriate symptomatic treatment be instituted immediately.
351

352

353 **5. PHARMACOLOGICAL PROPERTIES**

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355 **5.1 Pharmacodynamic properties**

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357 Pharmacotherapeutic group: Immunosuppressants, interleukin inhibitors, ATC code: L04AC13
358

359 Mechanism of action

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361 Ixekizumab is an IgG4 monoclonal antibody that binds with high affinity (< 3 pM) and specificity to
362 interleukin 17A (both IL-17A and IL-17A/F). Elevated concentrations of IL-17A have been implicated
363 in the pathogenesis of psoriasis by promoting keratinocyte proliferation and activation, as well as in

364 the pathogenesis of psoriatic arthritis and axial spondyloarthritis by driving inflammation leading to
365 erosive bone damage and pathological new bone formation. Neutralisation of IL-17A by ixekizumab
366 inhibits these actions. Ixekizumab does not bind to ligands IL-17B, IL-17C, IL-17D, IL-17E or
367 IL-17F.
368

369 In vitro binding assays confirmed that ixekizumab does not bind to human Fcγ receptors I, IIa, and IIIa
370 or to complement component C1q.

371 372 Pharmacodynamic effects

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374 Ixekizumab modulates biological responses that are induced or regulated by IL-17A. Based on
375 psoriatic skin biopsy data from a phase I study, there was a dose-related trend towards decreased
376 epidermal thickness, number of proliferating keratinocytes, T cells, and dendritic cells, as well as
377 reductions in local inflammatory markers from baseline to day 43. As a direct consequence treatment
378 with ixekizumab reduces erythema, induration and desquamation present in plaque psoriasis lesions.
379

380 Taltz has been shown to lower (within 1 week of treatment) levels of C-reactive protein, which is a
381 marker of inflammation.

382 383 Clinical efficacy and safety

384 385 *Adult plaque psoriasis*

386 The efficacy and safety of Taltz were assessed in three randomised, double-blind, placebo-controlled
387 phase III studies in adult patients (N=3,866) with moderate to severe plaque psoriasis who were
388 candidates for phototherapy or systemic therapy (UNCOVER-1, UNCOVER-2, and UNCOVER-3).
389 The efficacy and safety of Taltz were also evaluated versus etanercept (UNCOVER-2 and
390 UNCOVER-3). Patients randomised to Taltz who were sPGA (0,1) responders (static Physicians
391 Global Assessment) at week 12 were re-randomised to receive placebo or Taltz for an additional
392 48 weeks (UNCOVER-1 and UNCOVER-2); patients randomised to placebo, etanercept or Taltz who
393 were sPGA (0,1) non-responders received Taltz for up to 48 weeks. In addition, long term efficacy and
394 safety were evaluated in all three studies for up to a total of 5 years in patients who participated
395 through the entire study.
396

397 64 % of patients had received prior systemic therapy (biologic, conventional systemic or psoralen and
398 ultraviolet A (PUVA)), 43.5 % prior phototherapy, 49.3 % prior conventional systemic therapy, and
399 26.4 % prior biologic therapy 14.9 % had received at least one anti-TNF alpha agent, and 8.7 % an
400 anti-IL-12/IL-23. 23.4 % of patients had a history of psoriatic arthritis at baseline.
401

402 In all three studies, the co-primary endpoints were the proportion of patients who achieved a PASI 75
403 response (Psoriasis Area and Severity Index) and an sPGA of 0 (“clear”) or 1 (“minimal”) response at
404 week 12 versus placebo. The median baseline PASI score ranged from 17.4 to 18.3; 48.3 % to 51.2 %
405 of patients had a baseline sPGA score of severe or very severe, and mean baseline itch Numeric Rating
406 Scale (itch NRS) ranged from 6.3 to 7.1.
407

408 *Clinical response at 12 weeks*

409 UNCOVER-1 randomised 1,296 patients (1:1:1) to receive either placebo or Taltz (80 mg every two
410 or four weeks [Q2W or Q4W] following a 160 mg starting dose) for 12 weeks.
411

412 **Table 2. Efficacy results at week 12 in UNCOVER-1**
 413

Endpoints	Number of patients (%)			Difference from placebo in response rate (95% CI)	
	Placebo (N = 431)	Taltz 80 mg Q4W (N = 432)	Taltz 80 mg Q2W (N = 433)	Taltz 80 mg Q4W	Taltz 80 mg Q2W
sPGA of "0" (clear) or "1" (minimal)	14 (3.2)	330 (76.4) ^a	354 (81.8) ^a	73.1 (68.8, 77.5)	78.5 (74.5, 82.5)
sPGA of "0" (clear)	0	149 (34.5) ^a	160 (37.0) ^a	34.5 (30.0, 39.0)	37.0 (32.4, 41.5)
PASI 75	17 (3.9)	357 (82.6) ^a	386 (89.1) ^a	78.7 (74.7, 82.7)	85.2 (81.7, 88.7)
PASI 90	2 (0.5)	279 (64.6) ^a	307 (70.9) ^a	64.1 (59.6, 68.7)	70.4 (66.1, 74.8)
PASI 100	0	145 (33.6) ^a	153 (35.3) ^a	33.6 (29.1, 38.0)	35.3 (30.8, 39.8)
Itch NRS reduction $\geq 4^b$	58 (15.5)	305 (80.5) ^a	336 (85.9) ^a	65.0 (59.5, 70.4)	70.4 (65.4, 75.5)

414 *Abbreviations: N = number of patients in the intent-to-treat population*

415 *Note: patients with missing data were counted as non-responders*

416 ^a*p < 0.001 compared with placebo*

417 ^b*Patients with Itch NRS ≥ 4 at baseline: placebo N = 374, Taltz 80 mg Q4W N = 379, Taltz*
 418 *80 mg Q2W N = 391*

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 420 UNCOVER-2 randomised 1,224 patients (1:2:2:2) to receive either placebo, or Taltz (80 mg every two
 421 or four weeks [Q2W or Q4W] following a 160 mg starting dose) or etanercept 50 mg twice weekly for
 422 12 weeks.

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Table 3. Efficacy results at week 12 in UNCOVER-2

Endpoints	Number of patients (%)				Difference from placebo in response rate (95% CI)	
	Placebo (N = 168)	Taltz 80 mg Q4W (N = 347)	Taltz 80 mg Q2W (N = 351)	Etanercept 50 mg twice weekly (N = 358)	Taltz 80 mg Q4W	Taltz 80 mg Q2W
sPGA of “0” (clear) or “1” (minimal)	4 (2.4)	253 (72.9) ^{a,b}	292 (83.2) ^{a,b}	129 (36.0) ^a	70.5 (65.3, 75.7)	80.8 (76.3, 85.4)
sPGA of “0” (clear)	1 (0.6)	112 (32.3) ^{a,b}	147 (41.9) ^{a,b}	21 (5.9) ^c	31.7 (26.6, 36.7)	41.3 (36.0, 46.6)
PASI 75	4 (2.4)	269 (77.5) ^{a,b}	315 (89.7) ^{a,b}	149 (41.6) ^a	75.1 (70.2, 80.1)	87.4 (83.4, 91.3)
PASI 90	1 (0.6)	207 (59.7) ^{a,b}	248 (70.7) ^{a,b}	67 (18.7) ^a	59.1 (53.8, 64.4)	70.1 (65.2, 75.0)
PASI 100	1 (0.6)	107 (30.8) ^{a,b}	142 (40.5) ^{a,b}	19 (5.3) ^c	30.2 (25.2, 35.2)	39.9 (34.6, 45.1)
Itch NRS reduction $\geq 4^d$	19 (14.1)	225 (76.8) ^{a,b}	258 (85.1) ^{a,b}	177 (57.8) ^a	62.7 (55.1, 70.3)	71.1 (64.0, 78.2)

426 *Abbreviations: N = number of patients in the intent-to-treat population*

427 *Note: patients with missing data were counted as non-responders.*

428 ^a *p < 0.001 compared with placebo*

429 ^b *p < 0.001 compared with etanercept*

430 ^c *p < 0.01 compared with placebo*

431 ^d *Patients with Itch NRS ≥ 4 at baseline: placebo N = 135, Taltz 80 mg Q4W N = 293, Taltz*

432 *80 mg Q2W N = 303, etanercept N = 306*

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434 UNCOVER-3 randomised 1,346 patients (1:2:2:2) to receive either placebo, or Taltz (80 mg every two
435 or four weeks [Q2W or Q4W] following a 160 mg starting dose) or etanercept 50 mg twice weekly for
436 12 weeks.

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Table 4. Efficacy results at week 12 in UNCOVER-3

Endpoints	Number of patients (%)				Difference from placebo in response rate (95% CI)	
	Placebo (N = 193)	Taltz 80 mg Q4W (N = 386)	Taltz 80 mg Q2W (N = 385)	Etanercept 50 mg twice weekly (N = 382)	Taltz 80 mg Q4W	Taltz 80 mg Q2W
sPGA of “0” (clear) or “1” (minimal)	13 (6.7)	291 (75.4) ^{a,b}	310 (80.5) ^{a,b}	159 (41.6) ^a	68.7 (63.1, 74.2)	73.8 (68.5, 79.1)
sPGA of “0” (clear)	0	139 (36.0) ^{a,b}	155 (40.3) ^{a,b}	33 (8.6) ^a	36.0 (31.2, 40.8)	40.3 (35.4, 45.2)
PASI 75	14 (7.3)	325 (84.2) ^{a,b}	336 (87.3) ^{a,b}	204 (53.4) ^a	76.9 (71.8, 82.1)	80.0 (75.1, 85.0)
PASI 90	6 (3.1)	252 (65.3) ^{a,b}	262 (68.1) ^{a,b}	98 (25.7) ^a	62.2 (56.8, 67.5)	64.9 (59.7, 70.2)
PASI 100	0	135 (35.0) ^{a,b}	145 (37.7) ^{a,b}	28 (7.3) ^a	35 (30.2, 39.7)	37.7 (32.8, 42.5)
Itch NRS reduction $\geq 4^c$	33 (20.9)	250 (79.9) ^{a,b}	264 (82.5) ^{a,b}	200 (64.1) ^a	59.0 (51.2, 66.7)	61.6 (54.0, 69.2)

440 *Abbreviations: N = number of patients in the intent-to-treat population*

441 *Note: patients with missing data were counted as non-responders*

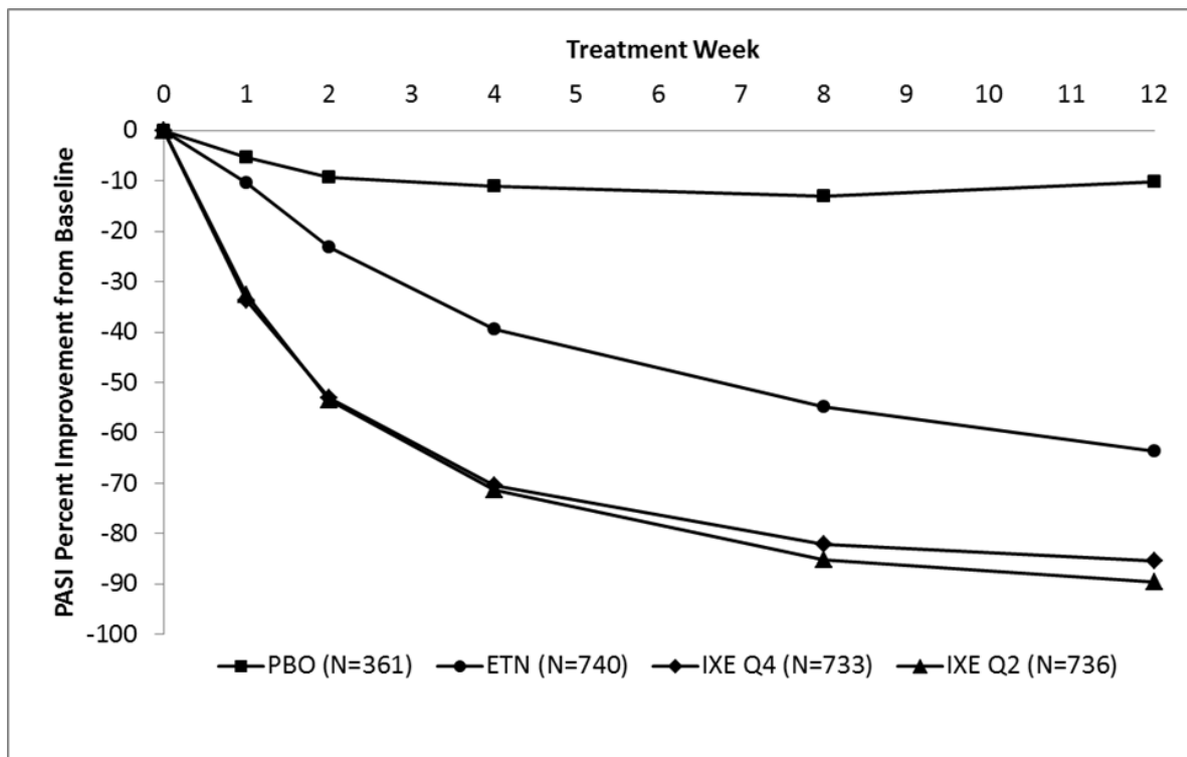
442 ^a *p < 0.001 compared with placebo*

443 ^b *p < 0.001 compared with etanercept*

444 ^c *Patients with Itch NRS ≥ 4 at baseline: placebo N = 158, Taltz 80 mg Q4W N = 313, Taltz*
445 *80 mg Q2W N = 320, etanercept N = 312*

446
447 Taltz was associated with a fast onset of efficacy with > 50 % reduction in mean PASI by week 2
448 (Figure 1). The percentage of patients achieving PASI 75 was significantly greater for Taltz compared
449 with placebo and etanercept as early as week 1. Approximately 25 % of patients treated with Taltz
450 achieved a PASI score < 5 by week 2, more than 55 % achieved the PASI score < 5 by week 4, and
451 increased to 85 % by Week 12 (compared to 3 %, 14 % and 50 % for etanercept). Significant
452 improvements in itch severity were seen at week 1 in patients treated with Taltz.
453

454 **Figure 1. PASI score, percent improvement at each post baseline visit (mBOCF) in the**
 455 **intent-to-treat population during the induction dosing period - UNCOVER-2 and**
 456 **UNCOVER-3**
 457



458
459

460 The efficacy and safety of Taltz was demonstrated regardless of age, gender, race, body weight, PASI
 461 baseline severity, plaques location, concurrent psoriatic arthritis, and previous treatment with a
 462 biologic. Taltz was efficacious in systemic treatment-naive, biologic-naive, biologic/anti-TNF-
 463 exposed and biologic/anti-TNF-failure patients.

464
 465 For patients identified as an sPGA (0,1) non-responder to etanercept at week 12 in UNCOVER-2
 466 (N = 200) and who were switched to Taltz 80 mg Q4W after a 4 week washout period, 73 % and
 467 83.5 % of patients achieved sPGA (0,1) and PASI 75, respectively, after 12 weeks of treatment with
 468 Taltz.

469
 470 In the 2 clinical studies that included an active comparator (UNCOVER-2 and UNCOVER-3), the rate
 471 of serious adverse events was 1.9 % for both etanercept and for Taltz, and the rate of discontinuation
 472 due to adverse events was 1.2 % for etanercept and 2.0 % for Taltz. The rate of infections was 21.5 %
 473 for etanercept and 26.0 % for Taltz, with 0.4 % being serious for etanercept and 0.5 % for Taltz.

474
 475 *Maintenance of response at week 60 and up to 5 years*
 476 Patients originally randomised to Taltz and who were responders at week 12 (i.e., sPGA score of 0,1)
 477 in UNCOVER-1 and UNCOVER-2 were re-randomised to an additional 48 weeks of treatment with
 478 placebo, or Taltz (80 mg every four or twelve weeks [Q4W or Q12W]).
 479 For sPGA (0,1) responders at week 12 re-randomised to treatment withdrawal (i.e., placebo), the
 480 median time to relapse (sPGA \geq 3) was 164 days in integrated UNCOVER- 1 and UNCOVER -2
 481 studies. Among these patients, 71.5 % regained at least an sPGA (0,1) response within 12 weeks of
 482 restarting treatment with Taltz 80 mg Q4W.

483

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Table 5. Maintenance of response and efficacy at week 60 (Studies UNCOVER-1 and UNCOVER-2)

Endpoints	Number of patients (%)				Difference from placebo in response rate (95% CI)	
	80 mg Q4W (induction) / Placebo (maintenance) (N = 191)	80 mg Q2W (induction) / Placebo (maintenance) (N = 211)	80 mg Q4W (induction) / 80 mg Q4W (maintenance) (N = 195)	80 mg Q2W (induction) / 80 mg Q4W (maintenance) (N = 221)	80 mg Q4W (induction) / 80 mg Q4W (maintenance)	80 mg Q2W (induction) / 80 mg Q4W (maintenance)
Maintained sPGA of “0” (clear) or “1” (minimal)	12 (6.3)	16 (7.6)	134 (68.7) ^a	173 (78.3) ^a	62.4 (55.1, 69.8)	70.7 (64.2, 77.2)
Maintained or achieved sPGA 0 (clear)	3 (1.6)	6 (2.8)	96 (49.2) ^a	130 (58.8) ^a	47.7 (40.4, 54.9)	56.0 (49.1, 62.8)
Maintained or achieved PASI 75	15 (7.9)	19 (9.0)	145 (74.4) ^a	184 (83.3) ^a	66.5 (59.3, 73.7)	74.3 (68.0, 80.5)
Maintained or achieved PASI 90	9 (4.7)	10 (4.7)	130 (66.7) ^a	169 (76.5) ^a	62.0 (54.7, 69.2)	71.7 (65.4, 78.0)
Maintained or achieved PASI 100	3 (1.6)	6 (2.8)	97 (49.7) ^a	127 (57.5) ^a	48.2 (40.9, 55.4)	54.6 (47.7, 61.5)

487 *Abbreviations: N = number of patients in the analysis population*

488 *Note: patients with missing data were counted as non-responders*

489 ^a *p < 0.001 compared with placebo*

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Taltz was efficacious in the maintenance of response in systemic treatment-naive, biologic-naive, biologic/anti-TNF-exposed and biologic/anti-TNF-failure patients.

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Significantly greater improvements at week 12 from baseline compared to placebo and etanercept were demonstrated in nail psoriasis (as measured by the Nail Psoriasis Severity Index [NAPSI]), in scalp psoriasis (as measured by Psoriasis Scalp Severity Index [PSSI]) and in palmoplantar psoriasis (as measured by Psoriasis Palmoplantar Severity Index [PPASI]) and were maintained at week 60 in patients treated with Taltz who were sPGA (0,1) responders at week 12.

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Of 591 subjects who received Taltz Q2W during the Induction Period then Q4W afterward in study UNCOVER 1, UNCOVER 2, and UNCOVER 3, 427 subjects completed 5 years of Taltz treatment, among those 101 patients required a dose escalation. Among the patients who completed the Week 264 assessment (N=427), 295 patients (69%), 289 patients (68%) and 205 patients (48%) were observed to have sPGA (0,1), PASI 90 and PASI 100 response, respectively, at Week 264. DLQI were collected after Induction Period in UNCOVER 1 and UNCOVER 2, 113 patients (66%) were observed to have DLQI (0,1) response.

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Quality of life/patient-reported outcomes

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At week 12 and across studies, Taltz was associated with statistically significant improvement in Health-related Quality of Life as assessed by mean decrease ranges from baseline in the Dermatology Life Quality Index (DLQI) (Taltz 80 mg Q2W from -10.2 to -11.1, Taltz 80 mg Q4W from -9.4 to -10.7, etanercept from -7.7 to -8.0 and placebo -1.0 to -2.0). A significantly greater proportion of patients treated with Taltz achieved a DLQI 0 or 1. Across studies a significantly greater proportion of patients treated with Taltz achieved a reduction of Itch NRS \geq 4 points at week 12 (84.6% for Taltz Q2W, 79.2% for Taltz Q4W and 16.5% for placebo) and the benefit was sustained over time up to

517 week 60 in patients treated with Taltz who were sPGA (0 or 1) responders at week 12. There was not
 518 any evidence of worsening of depression up to 60 weeks treatment with Taltz as assessed by the Quick
 519 Inventory of Depressive Symptomatology Self Report.

520
 521 *Postmarketing direct comparative studies*

522 IXORA-S: In a double-blind study ixekizumab Taltz was superior against ustekinumab on the primary
 523 study objective PASI 90 response at week 12 (Table 6). Onset of response was superior on PASI 75 as
 524 early as week 2 ($p < 0.001$) and on PASI 90 and PASI 100 by week 4 ($p < 0.001$). Superiority of Taltz
 525 versus ustekinumab was also demonstrated in the subgroups stratified by weight.

526
 527 **Table 6. PASI-response rates from comparative study ixekizumab versus ustekinumab**

	week 12		week 24		week 52	
	Taltz*	Ustekinumab**	Taltz*	Ustekinumab**	Taltz*	Ustekinumab**
Patients (n)	136	166	136	166	136	166
PASI 75, n (%)	120 (88.2 %)	114 (68.7 %)	124 (91.2 %)	136 (81.9%)	120 (88.2%)	126 (75.9 %)
PASI 90, n (%)	99 (72.8%) [§]	70 (42.2 %)	113 (83.1 %)	98 (59.0 %)	104 (76.5%)	98 (59.0 %)
PASI 100, n (%)	49 (36.0 %)	24 (14.5 %)	67 (49.3%)	39 (23.5 %)	71 (52.2%)	59 (35.5 %)

528 * Ixekizumab 160 mg given as a loading dose followed by 80 mg at week 2,4,6,8,10 and 12, and
 529 80 mg Q4W thereafter

530 ** Weight based dosing: Patients treated with ustekinumab received 45 mg or 90 mg at weeks 0 and
 531 4, then every 12 weeks until week 52 (dosed by weight as per approved posology)

532 [§] $p < 0.001$ versus ustekinumab (p value only provided for primary endpoint)

533
 534 IXORA R: Efficacy and safety of Taltz was also investigated in a 24 week randomized, double blind,
 535 parallel group study comparing Taltz to guselkumab, with Taltz being superior as early as Week 4 in
 536 achieving complete skin clearance and on the primary study objective (PASI 100 at week 12) and non
 537 inferior on PASI 100 at Week 24 (Table 7).

538
 539
 540 **Table 7. Efficacy Responses from comparative study ixekizumab versus guselkumab,**
 541 **Intent-to-Treat Population^a**

Endpoint	Time point	Guselkumab (N=507) response, n (%)	Ixekizumab (N=520) response, n (%)	Difference (IXE - GUS), % (CI)	p-value
Primary Objective					
PASI 100	Week 12	126 (24.9)	215 (41.3)	16.5 (10.8, 22.2)	<0.001
Major Secondary Objectives					
PASI 75	Week 2	26 (5.1)	119 (22.9)	17.8 (13.7, 21.8)	<0.001
PASI 90	Week 4	40 (7.9)	109 (21.0)	13.1 (8.9, 17.3)	<0.001
PASI 100	Week 4	7 (1.4)	35 (6.7)	5.4 (3.0, 7.7)	<0.001
PASI 90	Week 8	182 (35.9)	304 (58.5)	22.6 (16.6, 28.5)	<0.001
sPGA (0)	Week 12	128 (25.2)	218 (41.9)	16.7 (11.0, 22.4)	<0.001
PASI 50	Week 1	47 (9.3)	143 (27.5)	18.2 (13.6, 22.8)	<0.001
PASI 100	Week 8	69 (13.6)	154 (29.6)	16.0 (11.1, 20.9)	<0.001
PASI 100	Week 24	265 (52.3)	260 (50.0)	-2.3 (-8.4, 3.8)	0.414

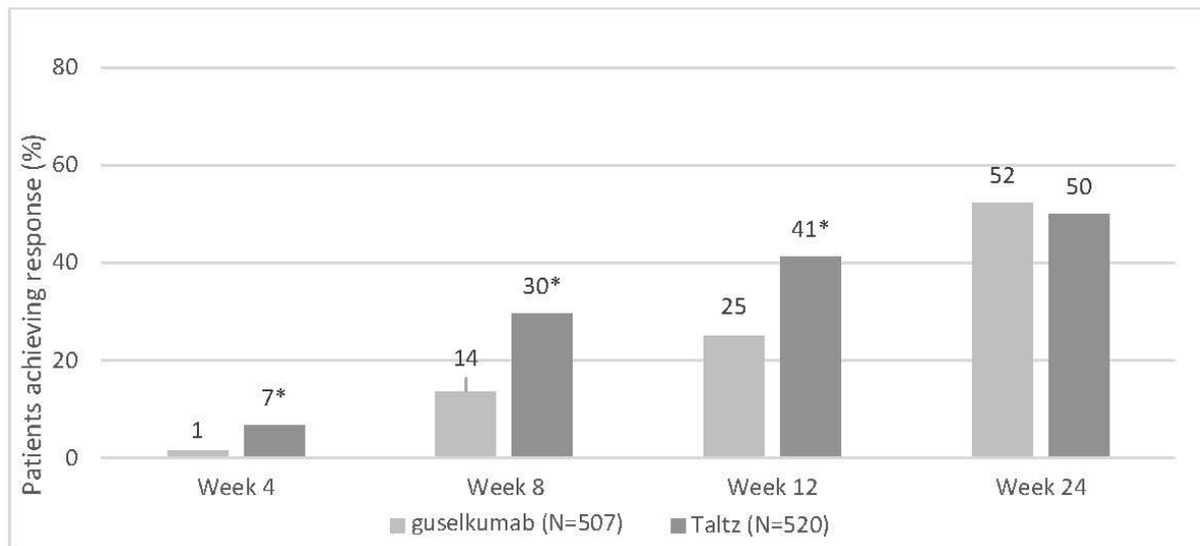
543 Abbreviations: CI = confidence interval; GUS = guselkumab; IXE = ixekizumab; N = number of
 544 patients in the analysis population; n = number of patients in the specified category; PASI =
 545 psoriasis area and severity index; sPGA = static physician global assessment.

546 ^a Endpoints were gated in this order

547
 548

549 **Figure 2: PASI 100 at weeks 4, 8, 12 and 24, NRI**

550



551

*p<0.001 vs guselkumab at weeks 4, 8, and 12

552

NRI = Non-Responder Imputation

553

554

555

556 *Efficacy in genital psoriasis*

557

558 A randomised, double-blind, placebo-controlled study (IXORA-Q) was conducted in 149 adult
559 subjects (24% females) with moderate to severe genital psoriasis (sPGA of Genitalia score of ≥ 3), a
560 minimum body surface area (BSA) involvement of 1% (60.4% had a BSA $\geq 10\%$) and previous failure
561 of or intolerance to at least one topical therapy for genital psoriasis. Patients had at least moderate
562 plaque psoriasis (defined as sPGA score of ≥ 3 and being candidates for phototherapy and/or systemic
563 therapy) for at least 6 months.

563

564 Subjects randomised to TALTZ received an initial dose of 160 mg followed by 80 mg every 2 weeks
565 for 12 weeks. The primary endpoint was the proportion of patients who achieved at least a "0" (clear)
566 or "1" (minimal) response on the sPGA of Genitalia (sPGA of Genitalia 0/1). At week 12, significantly
567 more subjects in the TALTZ group than placebo group achieved a sPGA of Genitalia 0/1 and a sPGA
568 0/1 independent of baseline BSA (baseline BSA 1% - <10% resp. $\geq 10\%$: sPGA of Genitalia "0" or
569 "1": Taltz 71%, resp. 75%; placebo: 0%, resp. 13%). A significantly greater proportion of patients
570 treated with TALTZ achieved a reduction in the PROs of severity of genital pain, genital itch, impact
571 of genital psoriasis on sexual activity, and Dermatology Quality of Life Index (DLQI).

572

573 **Table 8. Efficacy results at week 12 in Adults with genital psoriasis in trial IXORA-Q;**

574 **NRI^a**

575

Endpoints	TALTZ	Placebo	Difference from placebo (95% CI)
Number of patients (N) randomised	N=75	N=74	
sPGA of Genitalia "0" or "1"	73%	8%	65% (53%, 77%)
sPGA "0" or "1"	73%	3%	71% (60%, 81%)
DLQI 0,1 ^b	45%	3%	43% (31%, 55%)
N with baseline GPSS Itch NRS Score ≥ 3	N=62	N=60	
GPSS Genital Itch (≥ 3 point improvement)	60%	8%	51% (37%, 65%)
N with baseline SFQ Item 2 Score ≥ 2	N=37	N=42	
SFQ-item 2 score, "0" (never limited) or "1" (rarely limited)	78%	21%	57% (39%, 75%)

576

^a Abbreviations: NRI = Non-Responder Imputation; sPGA = static Physician Global Assessment;

577

GPSS = Genital Psoriasis Symptom Scale; SFQ = Sexual Frequency Questionnaire; DLQI =

578

Dermatology Quality of Life Index; ^b Total DLQI score of 0,1 indicates skin condition has no effect at

579 all on patient's life. sPGA of "0" or "1" is equivalent to "clear" or "minimal"; NRS = Numeric
 580 Rating Scale

581
 582 Paediatric plaque psoriasis
 583

584 A randomised, double-blind, multicenter, placebo-controlled trial (IXORA-Peds) enrolled
 585 201 children 6 to less than 18 years of age, with moderate to severe plaque psoriasis (as defined by a
 586 sPGA score ≥ 3 , involving $\geq 10\%$ of the body surface area, and a PASI score ≥ 12) who were candidates
 587 for phototherapy or systemic therapy, or were inadequately controlled on topical therapy.
 588 Patients were randomised to placebo (n=56), etanercept (n=30) or Taltz (n=115) with dosing stratified
 589 by weight:

- 590 <25 kg: 40 mg at week 0 followed by 20 mg Q4W (n=4)
- 591 25 kg to 50 kg: 80 mg at week 0 followed by 40 mg Q4W (n=50)
- 592 >50 kg: 160 mg at week 0 followed by 80 mg Q4W (n=147)

593 Patients randomized to etanercept (patients with severe psoriasis) received 0.8 mg/kg, not exceeding
 594 50 mg per dose, every week from week 0 through week 11.

595 Response to treatment was assessed after 12 weeks and defined by the proportion of patients who
 596 achieved the co-primary endpoint of an sPGA score of "0" (clear) or "1" (almost clear) with at least a
 597 2 point improvement from baseline and the proportion of patients that achieved a reduction in PASI
 598 score of at least 75% (PASI 75) from baseline.

599 Other evaluated outcomes at week 12 included the proportion of patients who achieved PASI 90,
 600 PASI 100, sPGA of "0" and an improvement of itch severity as measured by a reduction of at least
 601 4 points on an 11-point itch Numeric Rating Scale.

602 Patients had a median baseline PASI of 17 score ranging from 12-49. Baseline sPGA score was
 603 severe or very severe in 49%. Of all patients, 22% had received prior phototherapy and 32% had
 604 received prior conventional systemic therapy for the treatment of psoriasis.

605 25% of patients (n=43) were below 12 years (14% of patients [n=24] were 6-9 years and 11% of
 606 patients [n=19] were 10-11 years); 75% (n=128) were 12 years or above.

608 The clinical response data are presented in Table 9.

609
 610 **Table 9. Efficacy results in pediatric patients with plaque psoriasis, NRI**

Endpoints	Taltz ^a (N=115) n (%)	Placebo (N=56) n (%)	Difference vs placebo (95% CI)	Etanercept ^b (N=30) n (%)	Difference vs etanercept (95% CI) ^b
sPGA "0" (clear) or "1" (almost clear) ^c					
week 4	55 (48)	4 (7)	40.7 (29.3, 52.0) ^f	0(0)	36.8 (21.5, 52.2)
week 12 ^c	93 (81)	6 (11)	70.2 (59.3, 81.0) ^f	16 (53)	23.0 (0.6, 45.4)
sPGA "0" (clear) ^d	60 (52)	1 (2)	50.4 (40.6, 60.2) ^f	5 (17)	46.5 (26.2, 66.8)
PASI 75 ^c					
week 4	62 (54)	5 (9)	45.0 (33.2, 56.8) ^f	3 (10)	34.7 (15.6, 53.8)
week 12	102 (89)	14 (25)	63.7 (51.0, 76.4) ^f	19 (63)	20.9 (0.1, 41.7)
PASI 90 ^d	90 (78)	3 (5)	72.9 (63.3, 82.5) ^f	12 (40)	36.3 (14.2, 58.5)
PASI 100 ^d	57 (50)	1 (2)	47.8 (38.0, 57.6) ^f	5 (17)	43.9 (23.4, 64.3)
Itch NRS (≥ 4 point improvement) ^{d, e}	59 (71)	8 (20)	51.1 (35.3, 66.9) ^f	Not evaluated	---

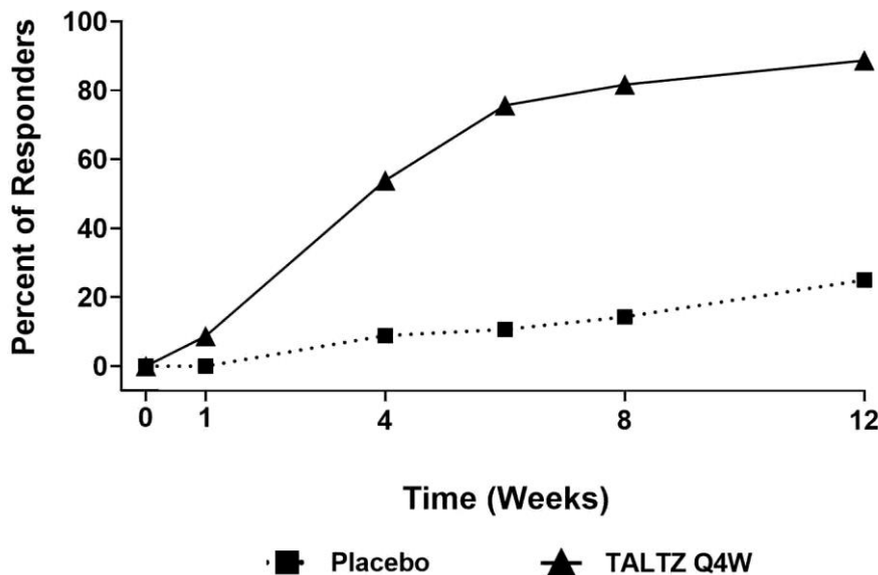
611 Abbreviations: N = Number of patients in the intent-to-treat population; NRI = Non-Responder
 612 Imputation.

613 ^a At week 0, subjects received 160 mg, 80 mg, or 40 mg of Taltz, followed by 80 mg, 40 mg, or
 614 20 mg every 4 weeks, depending on weight category, for 12 weeks.

615 ^b Comparisons to etanercept were performed within the sub-population of patients outside of US and
 616 Canada with severe Ps (N for Taltz = 38).

617 ^c Co-primary endpoints.

618 ^d Results at week 12.
 619 ^e Itch NRS (≥ 4 improvement) in patients with baseline Itch NRS ≥ 4 . The number of ITT patients
 620 with baseline Itch NRS Score ≥ 4 are as follows: Taltz, n = 83; PBO, n = 40.
 621 ^f p<0.001
 622
 623 **Figure 3. Percent of patients achieving PASI 75 in pediatric psoriasis through week 12**



624
 625
 626 Patients in the ixekizumab treatment group had clinically meaningful higher CDLQI/DLQI (0,1)
 627 responses at week 12 (NRI) compared with placebo. The difference between treatment groups was
 628 apparent from as early as week 4.
 629
 630 There were greater improvements at week 12 from baseline compared to placebo in nail psoriasis (as
 631 measured by the Nail Psoriasis Severity Index [NAPSI=0: Taltz 18% (6/34), placebo 0% (0/12)]), in
 632 scalp psoriasis (as measured by Psoriasis Scalp Severity Index [PSSI=0: Taltz 69% (70/102), placebo
 633 16% (8/50)]) and in palmoplantar psoriasis (as measured by Psoriasis Palmoplantar Severity Index
 634 [PPASI 75: Taltz 53% (9/17), placebo 11% (1/9)]).
 635
 636 Psoriatic arthritis
 637
 638 Taltz was assessed in two randomised, double-blind, placebo-controlled phase III studies in 780
 639 patients with active psoriatic arthritis (≥ 3 swollen and ≥ 3 tender joints). Patients had a diagnosis of
 640 psoriatic arthritis (Classification Criteria for Psoriatic Arthritis [CASPAR] criteria) for a median of 5.33
 641 years and had current plaque psoriasis skin lesions (94.0%) or a documented history of plaque
 642 psoriasis, with 12.1% of patients with moderate to severe plaque psoriasis at baseline. Over 58.9%
 643 and 22.3% of the psoriatic arthritis patients had enthesitis and dactylitis at baseline, respectively.
 644 Primary endpoint of both studies was American College of Rheumatology (ACR) 20 response at week
 645 24, followed by a long-term extension period from Week 24 to Week 156 (3 years).
 646
 647 In Psoriatic Arthritis Study 1 (SPIRIT-P1), patients naive to biologic therapy with active psoriatic
 648 arthritis were randomised to placebo, adalimumab 40 mg once every 2 weeks (active control reference
 649 arm), Taltz 80 mg once every 2 weeks (Q2W), or 80 mg once every 4 weeks (Q4W). Both Taltz
 650 regimens included a 160 mg starting dose. 85.3% of patients in this study had received prior treatment
 651 with ≥ 1 cDMARD. 53% of patients had concomitant use of MTX at a mean weekly dose of 15.8 mg.
 652 67% of patients who had concomitant use of MTX had a dose of 15 mg or greater. Patients with an
 653 inadequate response at week 16 received rescue therapy (modification to background therapy).
 654 Patients on Taltz Q2W or Q4W remained on their originally assigned dose of Taltz. Patients receiving

655 adalimumab or placebo were re-randomised 1:1 to Taltz Q2W or Q4W at week 16 or 24 based on
 656 responder status. 243 patients completed the extension period of 3 years on Taltz.
 657

658 Psoriatic Arthritis Study 2 (SPIRIT-P2) enrolled patients who were previously treated with an anti-TNF
 659 agent and discontinued the anti-TNF agent for either lack of efficacy or intolerance (anti-TNF-IR
 660 patients). Patients were randomised to placebo, Taltz 80 mg once every 2 weeks (Q2W), or 80 mg
 661 once every 4 weeks (Q4W). Both Taltz regimens included a 160 mg starting dose. 56% and 35% of
 662 patients were inadequate responders to 1 anti-TNF or 2 anti-TNF, respectively. SPIRIT-P2 evaluated
 663 363 patients, of whom 41% had concomitant use of MTX at a mean weekly dose of 16.1 mg. 73.2% of
 664 patients who had concomitant use of MTX had a dose of 15 mg or greater. Patients with an inadequate
 665 response at week 16 received rescue therapy (modification to background therapy). Patients in Taltz
 666 Q2W or Q4W remained on their originally assigned dose of Taltz. Patients receiving placebo were re-
 667 randomised 1:1 to Taltz Q2W or Q4W at week 16 or 24 based on responder status. 168 patients
 668 completed the extension period of 3 years on Taltz.

669 *Signs and symptoms*

670 Treatment with Taltz resulted in significant improvement in measures of disease activity compared to
 671 placebo at week 24 (see Table 10).
 672

673 **Table 10. Efficacy results in SPIRIT-P1 and SPIRIT-P2 at week 24**
 674

Endpoints	SPIRIT-P1					SPIRIT-P2					
	PBO (N = 106)	Taltz Q4W (N = 107)	Taltz Q2W (N = 103)	ADA (N = 101)	Difference from placebo in response rate (95% CI)		PBO (N = 118)	Taltz Q4W (N = 122)	Taltz Q2W (N = 123)	Difference from placebo in response rate (95% CI)	
ACR 20 response, n (%)											
week 24	32 (30.2)	62 (57.9)	64 (62.1)	58 (57.4)	27.8 (15.0, 40.6) ^c	31.9 (19.1, 44.8) ^c	23 (19.5)	65 (53.3)	59 (48.0)	33.8 (22.4, 45.2) ^c	28.5 (17.1, 39.8) ^c
ACR 50 response, n (%)											
week 24	16 (15.1)	43 (40.2)	48 (46.6)	39 (38.6)	25.1 (13.6, 36.6) ^c	31.5 (19.7, 43.3) ^c	6 (5.1)	43 (35.2)	41 (33.3)	30.2 (20.8, 39.5) ^c	28.3 (19.0, 37.5) ^c
ACR 70 response, n (%)											
week 24	6 (5.7)	25 (23.4)	35 (34.0)	26 (25.7)	17.7 (8.6, 26.8) ^c	28.3 (18.2, 38.5) ^c	0	27 (22.1)	15 (12.2)	22.1 (14.8, 29.5) ^c	12.2 (6.4, 18.0) ^c
Minimal disease activity (MDA) n (%)											
week 24	16 (15.1)	32 (29.9)	42 (40.8)	32 (31.7)	14.8 (3.8, 25.8) ^a	25.7 (14.0, 37.4) ^c	4 (3.4)	34 (27.9)	29 (23.6)	24.5 (15.9, 33.1) ^c	20.2 (12.0, 28.4) ^c
ACR 50 and PASI 100 in patients with ≥3% BSA psoriasis skin involvement at baseline, n (%)											
week 24	1 (1.5)	21 (28.8)	19 (32.2)	9 (13.2)	27.3 (16.5, 38.1) ^c	30.7 (18.4, 43.0) ^b	0 (0.0)	12 (17.6)	10 (14.7)	17.6 (8.6, 26.7) ^c	14.7 (6.3, 23.1) ^c

675 *Abbreviations: ACR 20/50/70 = American College of Rheumatology 20%/50%/70% response rate;*
 676 *ADA = adalimumab; BSA = body surface area; CI = confidence interval; Q4W = Taltz 80 mg every 4*
 677 *weeks; Q2W = Taltz 80 mg every 2 weeks; N = number of patients in the analysis population; n =*
 678 *number of patients in the specified category; NRI = non-responder imputation; PASI 100 = psoriasis*

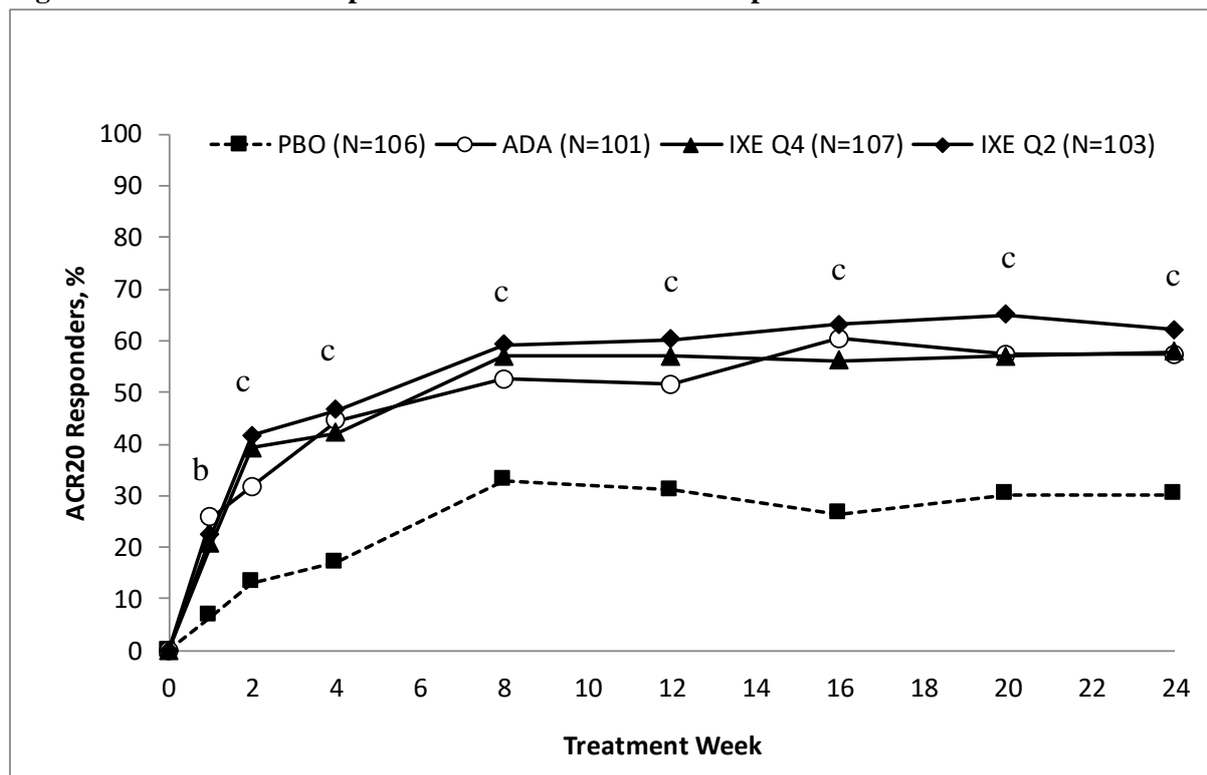
679 area and severity index 100% improvement; PBO = placebo.
 680 Note: patients who were rescued at week 16 or discontinued or with missing data were imputed as
 681 non-responders for week 24 analyses.
 682 Concomitant cDMARDs included MTX, leflunomide and sulfasalazine.
 683 a $p < 0.05$; b $p < 0.01$; c $p < 0.001$ compared with placebo.

684
 685 In patients with pre-existing dactylitis or enthesitis, treatment with Taltz Q4W resulted in
 686 improvement in dactylitis and enthesitis at week 24 compared to placebo (resolution: 78% vs. 24%;
 687 $p < 0.001$, and 39% vs. 21%; $p < 0.01$, respectively).
 688

689 In patients with $\geq 3\%$ BSA, the improvement in skin clearance at week 12 as measured by 75%
 690 improvement in Psoriasis Area Severity Index (PASI 75), was 67% (94/141) for those treated with the
 691 Q4W dosing regimen, and 9% (12/134) for those treated with placebo ($p < 0.001$). The proportion of
 692 patients achieving a PASI 75, PASI 90, and PASI 100 response at week 24 was greater with Taltz
 693 Q4W compared to placebo ($p < 0.001$). In patients with concomitant moderate to severe psoriasis and
 694 psoriatic arthritis, Taltz Q2W dose regimen showed significantly higher response rate for PASI75,
 695 PASI 90 and PASI 100 compared to placebo ($p < 0.001$) and demonstrated clinically meaningful benefit
 696 over the Q4W dose regimen.
 697

698 Treatment responses on Taltz were significantly greater than those on placebo as early as week 1 for
 699 ACR 20, week 4 for ACR 50 and week 8 for ACR 70 and persisted through week 24; effects were
 700 maintained through 3 years for patients who remained in the study.
 701

702 **Figure 4. ACR 20 response in SPIRIT-P1 over time up to week 24**



703
 704 For both Taltz Q2W and Q4W: b $p < 0.01$ and c $p < 0.001$ compared with placebo.
 705

706 In SPIRIT-P1 and SPIRIT-P2, similar responses for ACR 20/50/70 were seen in patients with psoriatic
 707 arthritis regardless of whether they were on concomitant cDMARDs, including MTX treatment, or
 708 not.
 709

710 In SPIRIT-P1 and SPIRIT-P2, improvements were shown in all components of the ACR scores
 711 including patient assessment of pain. At week 24 the proportion of patients achieving a modified
 712 Psoriatic Arthritis Response Criteria (PsARC) response was greater in the Taltz-treated patients
 713 compared to placebo.

714
715 In SPIRIT-P1, efficacy was maintained up to week 52 as assessed by ACR 20/50/70, MDA, enthesitis
716 resolution, dactylitis resolution, and PASI 75/90/100 response rates.

717
718 The efficacy and safety of Taltz was demonstrated regardless of age, gender, race, disease duration,
719 baseline body weight, baseline psoriasis involvement, baseline CRP, baseline DAS28-CRP,
720 concomitant corticosteroid use, and previous treatment with a biologic. Taltz was efficacious in
721 biologic-naive, biologic-exposed and biologic-failure patients.

722 In SPIRIT P1, 63 patients completed 3 years of Q4W ixekizumab treatment. Among the 107 patients
723 who were randomized to ixekizumab Q4W (NRI analysis in ITT population), 54 patients (50%), 41
724 patients (38%), 29 patients (27%), and 36 patients (34%) were observed to have ACR20, ACR50,
725 ACR70, and MDA response, respectively, at Week 156.

726 In SPIRIT P2, 70 patients completed 3 years of Q4W ixekizumab treatment. Among the 122 patients
727 who were randomized to ixekizumab Q4W (NRI analysis in ITT population), 56 patients (46%), 39
728 patients (32%), 24 patients (20%) and 33 (27%) were observed to have ACR20, ACR50, ACR70, and
729 MDA response, respectively, at Week 156.

730 *Radiographic response*

731 In SPIRIT-P1, inhibition of progression of structural damage was assessed radiographically and
732 expressed as the change in modified total Sharp Score (mTSS) and its components, the Erosion
733 Score (ES) and the Joint Space Narrowing score (JSN) at weeks 24 and 52, compared to baseline.
734 week 24 data are presented in Table 11.

735

736 **Table 11. Change in modified Total Sharp Score in SPIRIT-P1**

737

					Difference from placebo (95% CI)	
	PBO (N = 106)	Taltz Q4W (N = 107)	Taltz Q2W (N = 103)	ADA (N = 101)	Taltz Q4W	Taltz Q2W
Baseline score, mean (SD)	17.6 (28.62)	19.2 (32.68)	15.2 (28.86)	15.9 (27.37)	NA	NA
Change from baseline at week 24, LSM (SE)	0.51 (0.092)	0.18 (0.090)	0.09 (0.091)	0.13 (0.093)	-0.33 (-0.57,-0.09) ^b	-0.42 (-0.66,-0.19) ^c

738 *Abbreviations: ADA = adalimumab; CI = confidence interval; Q4W = Taltz 80 mg every 4 weeks;*
739 *Q2W = Taltz 80 mg every 2 weeks; LSM = least squares mean; N = number of patients in the analysis*
740 *population; PBO = placebo; SE = standard error; SD = standard deviation.*

741 *^b p<0.01; ^c p<0.001 compared with placebo.*

742

743 Radiographic joint damage progression was inhibited by Taltz (Table 11) at week 24, and the
744 percentage of patients with no radiographic joint damage progression (defined as a change from
745 baseline in mTSS of ≤ 0.5) from randomisation to week 24 was 94.8% for Taltz Q2W ($p < 0.001$), 89.0%
746 for Taltz Q4W ($p = 0.026$), 95.8% for adalimumab ($p < 0.001$), all compared to 77.4% for placebo. At
747 week 52, the mean change from baseline in mTSS was 0.27 for placebo/Taltz Q4W, 0.54 for Taltz
748 Q4W/Taltz Q4W, and 0.32 for adalimumab/Taltz Q4W. The percentage of patients with no
749 radiographic joint damage progression from randomisation to week 52 was 90.9% for placebo/Taltz
750 Q4W, 85.6% for Taltz Q4W/Taltz Q4W, and 89.4% for adalimumab/Taltz Q4W. Patients had no
751 structural progression from baseline (defined as $mTSS \leq 0.5$) in the treatment arms as follows:
752 Placebo/Taltz Q4W 81.5% (N=22/27), Taltz Q4W/Taltz Q4W 73.6% (N=53/72), and
753 adalimumab/Taltz Q4W 88.2% (N=30/34).

754 *Physical function and health-related quality of life*

755 In both SPIRIT-P1 and SPIRIT-P2, patients treated with Taltz Q2W ($p < 0.001$) and Q4W ($p < 0.001$)
756 showed significant improvement in physical function compared to patients treated with placebo as
757 assessed by Health Assessment Questionnaire-Disability Index (HAQ-DI) at week 24, and
758 maintained at week 52 in SPIRIT-P1.

759

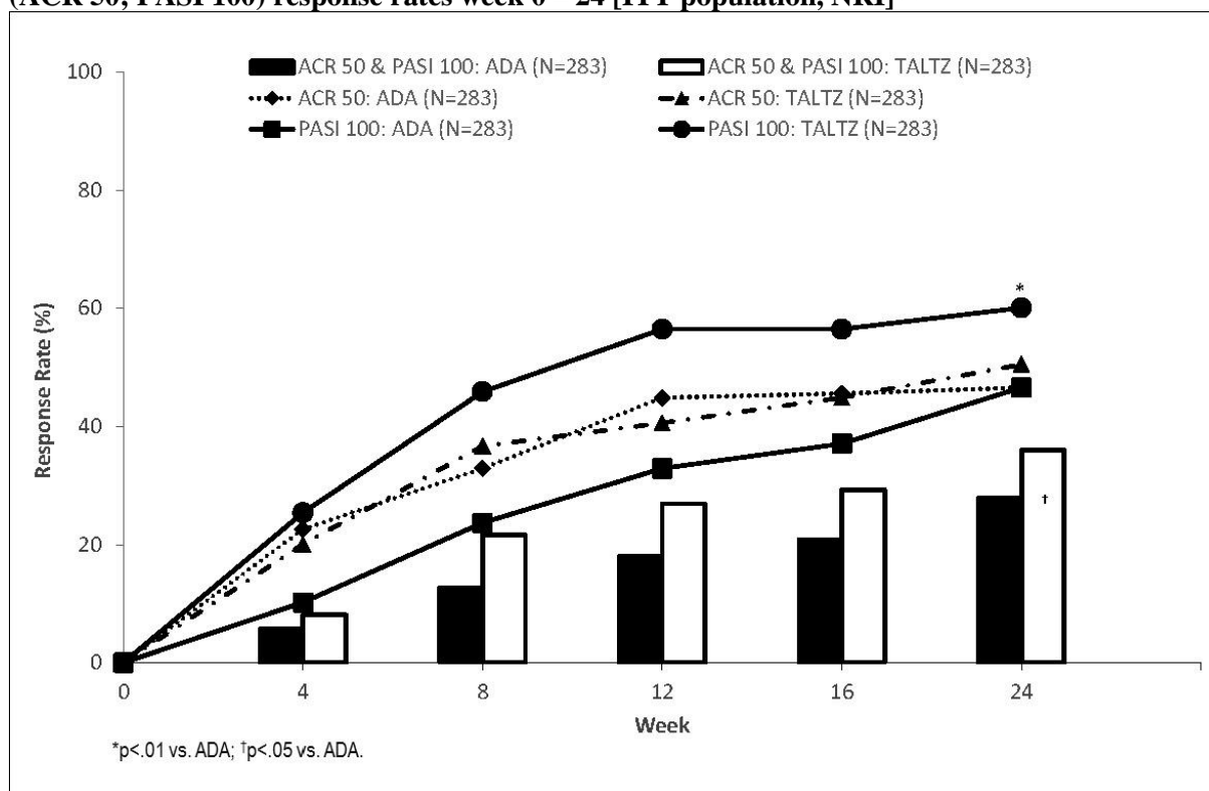
760 Taltz-treated patients reported improvements in health-related quality of life as measured by the
 761 Physical Component Summary of the Short Form-36 Health Survey (SF-36 PCS) score ($p < 0.001$).
 762 There were also improvements demonstrated in fatigue as assessed by Fatigue severity NRS scores
 763 ($p < 0.001$).
 764

765 Postmarketing phase 4, direct comparative study

766 Efficacy and safety of Taltz was investigated in a multicenter, randomised, open-label, rater-blinded,
 767 parallel-group study (SPIRIT-H2H) compared to adalimumab (ADA) in 566 patients with PsA who
 768 were naïve to biologic disease-modifying anti-rheumatic drugs (bDMARD). Patients were stratified
 769 at baseline based on concomitant cDMARD use and presence of moderate-to-severe psoriasis
 770 ($\text{PASI} \geq 12$, $\text{BSA} \geq 10$ and $\text{sPGA} \geq 3$).
 771

772 Taltz was superior to ADA on the primary study objective: simultaneous achievement of ACR 50
 773 and PASI 100 response at week 24 (Taltz 36.0% vs ADA 27.9%; $p = 0.036$; 95% confidence interval
 774 [0.5%, 15.8%]). Taltz also showed non-inferiority (pre-specified margin of -12%) to ADA on ACR
 775 50 (ITT analysis: Taltz 50.5% vs ADA 46.6%; 3.9% difference vs. ADA; 95% confidence interval [-
 776 4.3%; 12.1%]; PPS analysis Taltz: 52.3%, ADA: 53.1%, difference: -0.8% [CI: -10.3%; 8.7%]) and
 777 superiority on PASI 100 at week 24 (60.1% with Taltz vs 46.6% with ADA, $p = 0.001$), which were
 778 the major secondary endpoints in the study. At Week 52 a higher proportion of patients treated with
 779 Taltz versus ADA simultaneously achieved ACR50 and PASI 100 [39% (111/283) versus 26%
 780 (74/283)] and PASI 100 [64% (182/283) versus 41% (117/283)]. Taltz and ADA treatment resulted
 781 in similar responses for ACR50 [49.8% (141/283) versus 49.8% (141/283)]. Responses to Taltz
 782 were consistent when used as monotherapy or with concomitant use of methotrexate.
 783

784 **Figure 5: Primary endpoint (simultaneous ACR 50 & PASI 100) and major secondary endpoints**
 785 **(ACR 50; PASI 100) response rates week 0 – 24 [ITT population, NRI]****



786 ** Taltz 160 mg week 0, then 80 mg every 2 weeks to week 12 and every 4 weeks thereafter for
 787 patients with moderate to severe plaque psoriasis or 160 mg week 0, then 80 mg every 4 week for
 788 other patients, ADA 80 mg week 0, then 40 mg every 2 weeks from week 1 for patients with moderate
 789 to severe plaque psoriasis or 40 mg week 0, then 40 mg every 2 weeks for other patients.
 790 Significance level only provided for endpoint that was pre-defined and multiplicity tested.
 791

792 Axial spondyloarthritis

793
 794

795 Taltz was assessed in a total of 960 adult patients with axial spondyloarthritis in three randomised
 796 placebo-controlled studies (two in radiographic and one in non-radiographic axial spondyloarthritis).
 797

798 Radiographic axial spondyloarthritis

799 Taltz was assessed in a total of 657 patients in two randomised, double-blind, placebo-controlled
 800 studies (COAST-V and COAST-W) in adult patients who had active disease as defined by the Bath
 801 Ankylosing Spondylitis Disease Activity Index (BASDAI) ≥ 4 and total back pain ≥ 4 on a numeric
 802 rating scale despite non-steroidal anti-inflammatory drug (NSAID) therapy. Across both studies at
 803 baseline, patients had symptoms for a mean of 17 years (median of 16 years). At baseline,
 804 approximately 32% of the patients were on a concomitant cDMARD.

805 COAST-V evaluated 341 biologic-naive patients treated with either Taltz 80 mg or 160 mg at week 0
 806 followed by 80 mg every 2 weeks (Q2W) or 4 weeks (Q4W), adalimumab 40 mg every 2 weeks, or
 807 with placebo. Patients receiving placebo were re-randomised at week 16 to receive Taltz (160 mg
 808 starting dose, followed by 80 mg Q2W or Q4W). Patients receiving adalimumab were re-randomised
 809 at week 16 to receive Taltz (80 mg Q2W or Q4W).

810 COAST-W evaluated 316 patients who had prior experience with 1 or 2 TNF-inhibitors (90% were
 811 inadequate responders and 10% were intolerant to TNF inhibitors). All patients were treated with
 812 Taltz 80 or 160 mg at week 0 followed by 80 mg Q2W or Q4W, or with placebo. Patients receiving
 813 placebo were re-randomised at week 16 to receive Taltz (160 mg initial dose, followed by 80 mg Q2W
 814 or Q4W).

815 The primary endpoint in both studies was the percentage of patients achieving an Assessment of
 816 Spondyloarthritis International Society 40 (ASAS40) response at week 16.

817

818 *Clinical response*

819 In both studies, patients treated with Taltz 80 mg Q2W or 80 mg Q4W demonstrated greater
 820 improvements in ASAS40 and ASAS20 responses compared to placebo at week 16 (Table 12).

821 Responses were similar in patients regardless of concomitant therapies. In COAST-W, responses were
 822 seen regardless of the number of prior TNF inhibitors.

823

824 **Table 12. Efficacy results in COAST-V and COAST-W at week 16**

825

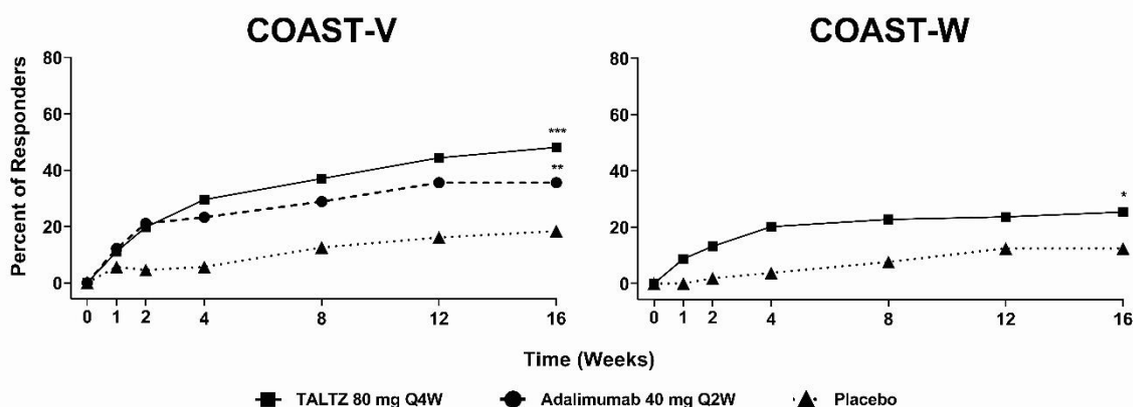
	COAST-V, biologic-naive				COAST-W, TNF-inhibitor experienced		
	Taltz 80 mg Q4W ^a (N=81)	Placebo (N=87)	Difference from placebo ^g	Adalimumab 40 mg Q2W (N=90)	Taltz 80 mg Q4W ^c (N=114)	Placebo (N=104)	Difference from placebo ^g
ASAS20 response ^b , n (%), NRI	52 (64.2%)	35 (40.2%)	24.0 (9.3, 38.6)**	53 (58.9%)	55 (48.2%)	31 (29.8%)	18.4 (5.7, 31.1) **
ASAS40 response ^{b,c} , n (%), NRI	39 (48.1%)	16 (18.4%)	29.8 (16.2, 43.3)***	32 (35.6%)	29 (25.4%)	13 (12.5%)	12.9 (2.7, 23.2) *
ASDAS							
Change from baseline <i>Baseline</i>	-1.4 3.7	-0.5 3.9	-1.0 (-1.3, -0.7) ***	-1.3*** 3.7	-1.2 4.2	-0.1 4.1	-1.1 (-1.3, -0.8) ***
BASDAI Score							
Change from baseline <i>Baseline</i>	-2.9 6.8 ⁱ	-1.4 6.8 ⁱ	-1.5 (-2.1, -0.9) ***	-2.5*** 6.7 ⁱ	-2.2 7.5	-0.9 7.3	-1.2 (-1.8, -0.7) ***
MRI Spine SPARCC^d							
Change from baseline <i>Baseline</i>	-11.0 14.5	-1.5 15.8	-9.5 (-12.6, - 6.4)***	-11.6*** 20.0	-3.0 8.3	3.3 6.4	-6.3 (-10.0, - 2.5)**
BASDAI50 ^e n (%), NRI	34 (42.0%)	15 (17.2%)	24.7 (11.4, 38.1)***	29 (32.2%)*	25 (21.9%) ⁱ	10 (9.6%) ⁱ	12.3 (2.8, 21.8)*
ASDAS <2.1, n (%) (low disease activity), NRI	35 (43.2%) ^h	11 (12.6%) ^h	30.6 (17.7, 43.4)***	34 (37.8%)*** ^h	20 (17.5%)	5 (4.8%)	12.7 (4.6, 20.8)**
ASDAS <1.3, n (%) (inactive disease), NRI	13 (16.0%)	2 (2.3%)	13.8 (5.2, 22.3) **	14 (15.6%)* ^h	4 (3.5%) ⁱ	1 (1.0%) ⁱ	2.5 (-1.3, 6.4)
ASAS HI ^f Change from baseline <i>Baseline</i>	-2.4 7.5	-1.3 8.1	-1.1 (-2.0, -0.3) *	-2.3* 8.2	-1.9 10.0	-0.9 9.0	-1.0 (-1.9, -0.1) *
SF-36 PCS Change from baseline <i>Baseline</i>	7.7 34.0	3.6 32.0	4.1 (1.9, 6.2) ***	6.9** 33.5	6.6 27.5	1.4 30.6	5.2 (3.0, 7.4) ***

826 Abbreviations: N = number of patients in the intent-to-treat population; NRI = Non-responder
 827 Imputation; patients with missing data were counted as non-responders.
 828 ASAS HI = Assessment of SpondyloArthritis International Society Health Index; ASDAS = Ankylosing
 829 Spondylitis Disease Activity Score; BASDAI = Bath Ankylosing Spondylitis Disease Activity Index;
 830 CFB = least square mean change from baseline at week 16; MRI Spine SPARCC = Spondyloarthritis
 831 Research Consortium of Canada Magnetic Resonance Imaging Scoring of the Spine
 832 (23 discovertebral unit scale)

- 833
 834 ^a At week 0, patients received 80 mg or 160 mg of Taltz.
 835 ^b An ASAS20 response is defined as a $\geq 20\%$ improvement and an absolute improvement from
 836 baseline of ≥ 1 unit (range 0 to 10) in ≥ 3 of 4 domains (Patient Global, Spinal Pain, Function, and
 837 Inflammation), and no worsening of $\geq 20\%$ and ≥ 1 unit (range 0 to 10) in the remaining domain. An
 838 ASAS40 response is defined as a $\geq 40\%$ improvement and an absolute improvement from baseline
 839 of ≥ 2 units in ≥ 3 of 4 domains without any worsening in the remaining domain.
 840 ^c Primary endpoint.
 841 ^d The numbers of ITT patients with MRI data at baseline are as follows: COAST-V: Taltz, n = 81;
 842 PBO, n = 82; ADA, n=85. COAST-W: Taltz, n = 58; PBO, n = 51.
 843 ^e BASDAI50 response defined as an improvement of $\geq 50\%$ of the BASDAI score from baseline.
 844 ^f ASAS HI: Assessment of SpondyloArthritis International Society Health Index (ASAS HI) across all
 845 domains.
 846 ^g The reported values are difference in % (95% CI) for categorical variables, and difference in
 847 LSM(95% CI) for continuous variables.
 848 ^h post hoc analysis, not multiplicity corrected.
 849 ⁱ prespecified, but not multiplicity gated.
 850 * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$ compared with placebo.

851
 852 There were improvements in the main components of the ASAS40 response criteria (spinal pain,
 853 BASFI, patient global assessment, stiffness) and other measures of disease activity, including CRP, at
 854 week 16.

855
 856
 857
 858 **Figure 6. Percent of patients achieving ASAS40 responses in COAST-V and COAST-W**
 859 **through week 16, NRI^a**



- 861
 862
 863 ^a Patients with missing data were counted as non-responders.
 864 * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$ compared with placebo.

865
 866 Similar response in ASAS40 was seen in patients regardless of baseline CRP levels, baseline ASDAS
 867 scores and MRI spine SPARCC scores. The ASAS40 response was demonstrated regardless of age,
 868 gender, race, disease duration, baseline body weight, baseline BASDAI score and prior biologic
 869 treatment.

870 In COAST-V and COAST-W efficacy was maintained up to week 52 as assessed by the endpoints
 871 presented in Table 12, including ASAS20, ASAS40, ASDAS, BASDAI, and ASAS HI response rates.

872
 873 *Health-related outcomes*

874 Spinal pain showed improvements versus placebo as early as week 1, maintained through week 16
 875 [Taltz vs placebo: COAST-V -3.2 vs -1.7; COAST-W -2.4 vs -1.0]; fatigue and spinal mobility
 876 showed improvements versus placebo at week 16. Improvements in spinal pain, fatigue and spinal
 877 mobility were maintained through week 52.

878
 879 *Non-radiographic axial spondyloarthritis*

880 Taltz was assessed in a randomised, double-blind, study with a 52-week placebo-controlled period
 881 (COAST-X) in 303 adult patients with active axial spondyloarthritis for at least 3 months. Patients
 882 must have had objective signs of inflammation indicated by elevated C-reactive protein (CRP) and/or
 883 sacroiliitis on magnetic resonance imaging (MRI), and no definitive radiographic evidence of
 884 structural damage on sacroiliac joints. Patients had active disease as defined by the Bath Ankylosing
 885 Spondylitis Disease Activity Index (BASDAI) ≥ 4 , and spinal pain ≥ 4 on a 0 to 10 Numerical Rating
 886 Scale (NRS), despite non-steroidal anti-inflammatory drug (NSAID) therapy. Patients were treated
 887 with either Taltz 80 mg or 160 mg at week 0, followed by 80 mg every 2 weeks (Q2W) or 80 mg
 888 every 4 weeks (Q4W) or with placebo. Dose adjustment and/or initiation of concomitant medications
 889 (NSAIDs, cDMARDs, corticosteroids, analgesics) were permitted starting at week 16.

890
 891 At baseline, patients had symptoms of non-radiographic axSpA for an average of 11 years.
 892 Approximately 39% of the patients were on a concomitant cDMARD.

893
 894 The primary endpoint was the percentage of patients achieving an Assessment of Spondyloarthritis
 895 International Society 40 (ASAS40) response at week 16.

896
 897 *Clinical response*

898 Higher proportions of patients treated with Taltz 80 mg Q4W achieved ASAS40 response compared to
 899 placebo at week 16 (Table 13). Responses were similar regardless of concomitant therapies.

900
 901
 902

Table 13. Efficacy results at week 16 in COAST-X, NRI ^{a,b}

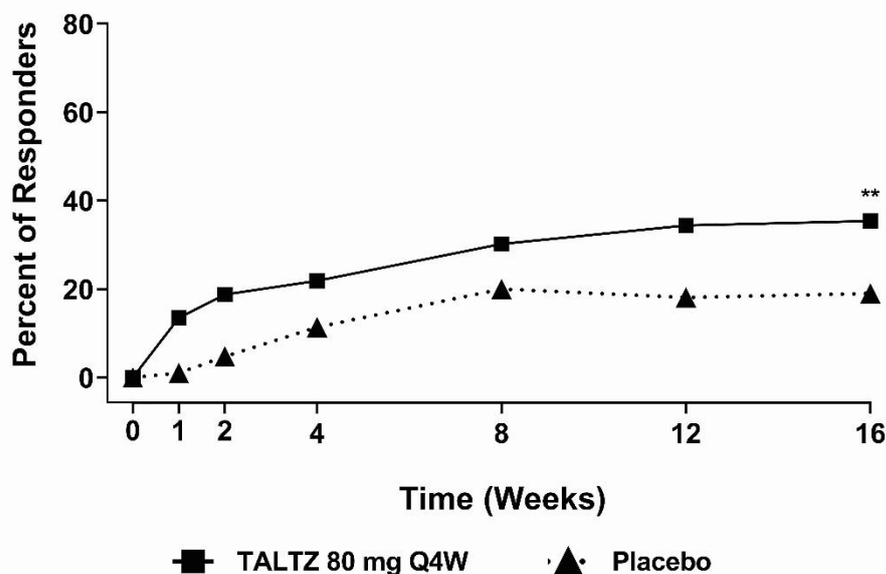
	Taltz 80 mg Q4W^c (N=96)	Placebo (N=105)	Difference from placebo ^h
ASAS20 response ^d , n (%), NRI	52 (54.2%)	41 (39.0%)	15.1 (1.5, 28.8)*
ASAS40 response ^{d,e} , n (%), NRI	34 (35.4%)	20 (19.0%)	16.4 (4.2, 28.5)**
ASDAS			
Change from baseline	-1.1	-0.6	-0.5 (-0.8, -0.3) ***
<i>Baseline</i>	3.8	3.8	
BASDAI Score			
Change from baseline	-2.2	-1.5	-0.7 (-1.3, -0.1) *
<i>Baseline</i>	7.0	7.2	
MRI SIJ SPARCC^f			
Change from baseline	-3.4	-0.3	-3.1 (-4.6, -1.6) ***
<i>Baseline</i>	5.1	6.3	
ASDAS <2.1, n (%) (low disease activity), NRI ^g	26 (27.7%)	13 (12.4%)	15.3 (4.3, 26.3) **
SF-36 PCS			
Change from baseline	8.1	5.2	2.9 (0.6, 5.1) *
<i>Baseline</i>	33.5	32.6	

903 ^a Abbreviations: N = number of patients in the intent-to-treat population; NRI = Non-responder
 904 Imputation. ASDAS = Ankylosing Spondylitis Disease Activity Score; BASDAI = Bath Ankylosing
 905 Spondylitis Disease Activity Index; Change from baseline = least square mean change from baseline
 906 at week 16; MRI SIJ SPARCC = Spondyloarthritis Research Consortium of Canada Magnetic
 907 Resonance Imaging Scoring of the sacroiliac joint.

908 ^b Patients with missing data were counted as non-responders.
 909 ^c At week 0, patients received 80 mg or 160 mg of Taltz.
 910 ^d An ASAS20 response is defined as a $\geq 20\%$ improvement and an absolute improvement from baseline
 911 of ≥ 1 units (range 0 to 10) in ≥ 3 of 4 domains (Patient Global, Spinal Pain, Function, and
 912 Inflammation), and no worsening of $\geq 20\%$ and ≥ 1 unit (range 0 to 10) in the remaining domain. An
 913 ASAS40 response is defined as a $\geq 40\%$ improvement and an absolute improvement from baseline of
 914 ≥ 2 units in ≥ 3 of 4 domains without any worsening in the remaining domain.
 915 ^e Primary endpoint at week 16.
 916 ^f The numbers of ITT patients with MRI data at baseline and week 16 are as follows: Taltz, $n = 85$;
 917 PBO, $n = 90$.
 918 ^g Patients with missing data were counted as non-responders. Percentages are based on the number of
 919 patients in the ITT population with baseline ASDAS ≥ 2.1 .
 920 ^h The reported values are difference in % (95% CI) for categorical variables, and difference in
 921 LSM (95% CI) for continuous variables.
 922 * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$ compared with placebo.

923
 924 The improvement in the main components of the ASAS40 response criteria (spinal pain, BASFI,
 925 patient global assessment, stiffness) and other measures of disease activity demonstrated significant
 926 clinical improvement at week 16.
 927
 928

929 **Figure 7. Percent of patients achieving ASAS40 response through week 16 in COAST-X,**
 930 **NRI^a**



931
 932 ^a Patients with missing data were counted as non-responders.
 933 ** $p < 0.01$ compared with placebo.
 934

935 Efficacy was maintained up to week 52 as assessed by the endpoints presented in Table 13.

936 Health-related outcomes

937 Spinal pain showed improvements versus placebo as early as week 1 and was maintained through
 938 week 16 [Taltz vs placebo: COAST-X: -2.4 vs -1.5]. In addition, more patients on Taltz compared
 939 with placebo achieved good health status (ASAS HI ≤ 5) at week 16 and week 52.
 940

941 Immunisations

942 In a study in healthy subjects, no safety concerns were identified of two inactivated vaccines (tetanus
 943 and pneumococcal), received after two doses of ixekizumab (160 mg followed by a second dose of 80
 944

945 mg two weeks later). However, the data concerning immunisation were insufficient to conclude on an
946 adequate immune response to these vaccines following administration of Taltz.

947

948 Paediatric population

949

950 The European Medicines Agency has deferred the obligation to submit the results of studies with Taltz
951 in one or more subsets of the paediatric population in the treatment of plaque psoriasis and psoriatic
952 arthritis/axial spondyloarthritis (see section 4.2 for information on paediatric use).

953

954 **5.2 Pharmacokinetic properties**

955

956 Absorption

957

958 Following a single subcutaneous dose of ixekizumab in patients with psoriasis, mean peak
959 concentrations were achieved within 4 to 7 days, across a dose range of 5 to 160 mg. The mean (SD)
960 maximum plasma concentration (C_{max}) of ixekizumab, after the 160 mg starting dose, was
961 19.9 (8.15) $\mu\text{g/ml}$.

962

963 After the 160 mg starting dose, steady state was achieved by week 8 with the 80 mg Q2W dosing
964 regimen. Mean (SD) $C_{max,ss}$, and $C_{trough,ss}$ estimates are 21.5 (9.16) $\mu\text{g/ml}$, and 5.23 (3.19) $\mu\text{g/ml}$.

965

966 After switching from the 80 mg Q2W dosing regimen to the 80 mg Q4W dosing regimen at week 12,
967 steady state would be achieved after approximately 10 weeks. Mean (SD) $C_{max,ss}$, and $C_{trough,ss}$ estimates
968 are 14.6 (6.04) $\mu\text{g/ml}$, and 1.87 (1.30) $\mu\text{g/ml}$.

969

970 The average bioavailability of ixekizumab after subcutaneous administration was 54 % to 90 % across
971 analyses.

972

973 Distribution

974

975 From population pharmacokinetic analyses, the mean total volume of distribution at steady state was
976 7.11 L.

977

978 Biotransformation

979

980 Ixekizumab is a monoclonal antibody and is expected to be degraded into small peptides and amino
981 acids via catabolic pathways in the same manner as endogenous immunoglobulins.

982

983 Elimination

984

985 In the population PK analysis, mean serum clearance was 0.0161 L/hr. Clearance is independent of
986 dose. The mean elimination half-life, as estimated from population pharmacokinetic analysis, is
987 13 days in patients with plaque psoriasis.

988

989 Linearity/non-linearity

990

991 Exposure (AUC) increased proportionally over a dose range of 5 to 160 mg given as a subcutaneous
992 injection.

993

994

995 Pharmacokinetic properties across indications

996

997 The pharmacokinetic properties of Taltz were similar across the plaque psoriasis, psoriatic arthritis,
998 radiographic axial spondyloarthritis and non-radiographic axial spondyloarthritis indications.

999

1000

1001 Elderly

1002
1003 Of the 4,204 plaque psoriasis patients exposed to Taltz in clinical studies, a total of 301 were 65 years
1004 of age or older and 36 patients were 75 years of age or older. Of the 1,118 psoriatic arthritis patients
1005 exposed to Taltz in clinical studies, a total of 122 patients were 65 years of age or older and 6 patients
1006 were 75 years of age or older.

1007 Based on population pharmacokinetic analysis with a limited number of elderly patients (n = 94 for
1008 age ≥ 65 years and n = 12 for age ≥ 75 years), clearance in elderly patients and patients less than
1009 65 years of age was similar.

1010

1011 Renal or hepatic impairment

1012

1013 Specific clinical pharmacology studies to evaluate the effects of renal impairment and hepatic
1014 impairment on the PK of ixekizumab have not been conducted. Renal elimination of intact
1015 ixekizumab, an IgG MAb, is expected to be low and of minor importance; similarly, IgG MAbs are
1016 mainly eliminated via intracellular catabolism and hepatic impairment is not expected to influence
1017 clearance of ixekizumab.

1018

1019 Paediatric population

1020

1021 Paediatric psoriasis patients (age 6 to less than 18 years) were administered ixekizumab at the
1022 recommended paediatric dosing regimen for 12 weeks. Patients weighing >50 kg and 25 to 50 kg had
1023 a mean ±SD steady-state trough concentration of 3.8 ±2.2 µg/ml and 3.9 ±2.4 µg/ml, respectively, at
1024 week 12.

1025

1026 **5.3 Preclinical safety data**

1027

1028 Non-clinical data reveal no special hazards for humans based on repeat-dose toxicity studies, safety
1029 pharmacology evaluations, and reproductive and developmental toxicity studies.

1030

1031 Ixekizumab administration to cynomolgus monkeys for 39 weeks at subcutaneous doses up to
1032 50 mg/kg weekly produced no organ toxicity or undesirable effects on immune function (e.g. T-cell
1033 dependent antibody response and NK cell activity). A weekly subcutaneous dose of 50 mg/kg to
1034 monkeys is approximately 19 times the 160 mg starting dose of Taltz and in monkeys results in
1035 exposure (AUC) that is at least 61-fold higher than the predicted mean steady-state exposure in
1036 humans administered the recommended dose regimen.

1037

1038 Non-clinical studies have not been conducted to evaluate the carcinogenic or mutagenic potential of
1039 ixekizumab.

1040

1041 No effects on reproductive organs, menstrual cycles or sperm were observed in sexually mature
1042 cynomolgus monkeys that received ixekizumab for 13 weeks at a weekly subcutaneous dose of
1043 50 mg/kg.

1044

1045 In developmental toxicity studies, ixekizumab was shown to cross the placenta and was present in the
1046 blood of offspring for up to 6 months of age. A higher incidence of postnatal mortality occurred in the
1047 offspring of monkeys given ixekizumab compared to concurrent controls. This was related primarily
1048 to early delivery or maternal neglect of offspring, common findings in nonhuman primate studies, and
1049 considered clinically irrelevant.

1050

1051 **6. PHARMACEUTICAL PARTICULARS**

1052

1053 **6.1 List of excipients**

1054

1055 Sodium citrate

1056 Citric acid, anhydrous

1057 Sodium chloride

1058 Polysorbate 80
1059 Water for injections

1060

1061 **6.2 Incompatibilities**

1062

1063 Not applicable.

1064

1065 **6.3 Shelf life**

1066

1067 2 years.

1068

1069 **6.4 Special precautions for storage**

1070

1071 Store in a refrigerator (2 °C to 8 °C).

1072 Do not freeze.

1073 Store in the original package in order to protect from light.

1074

1075 Taltz may be stored unrefrigerated for up to 5 days at a temperature not above 30 °C.

1076

1077 **6.5 Nature and contents of container**

1078

1079 1 ml solution in a type I clear glass syringe.

1080 Pack sizes of 1, 2, or 3 pre-filled syringes.

1081 Not all pack sizes may be marketed.

1082

1083

1084 **6.6 Special precautions for disposal and other handling**

1085

1086 The instructions for using the syringe, included with the package leaflet, must be followed carefully.

1087

1088 The pre-filled syringe is for single use only.

1089

1090 Taltz should not be used if particles appear or if the solution is cloudy and/or distinctly brown.

1091

1092 Taltz that has been frozen must not be used.

1093

1094 Any unused medicinal product or waste material should be disposed of in accordance with local requirements.

1095

1096

1097 40 mg preparation of ixekizumab for children 25-50 kg body weight

1098 Ixekizumab doses of 40 mg must be prepared and administered by a qualified healthcare professional.

1099 Use only Taltz 80 mg solution for injection prefilled-syringe when preparing the prescribed 40 mg paediatric doses.

1100

1101

- 1102 1. Expel the entire contents of the prefilled syringe into a sterile, clear glass vial. DO NOT
- 1103 shake or swirl the vial.
- 1104 2. Use a 0.5 mL or 1 mL disposable syringe and sterile needle to withdraw the prescribed dose
- 1105 (0.5 ml for 40 mg) from the vial.
- 1106 3. Change the needle and use a 27-gauge, sterile needle to inject the patient. Discard any
- 1107 unused ixekizumab in the vial.

1108

1109 The prepared ixekizumab must be administered within 4 hours of puncturing the sterile vial at room temperature.

1110

1111

1112 **7. MARKETING AUTHORISATION HOLDER**

1113

1114 Zuellig Pharma Ltd. Bangkok, Thailand

1115

1116 **8. MARKETING AUTHORISATION NUMBER(S)**

1117

1118 1C 15084/63 (NBC)

1119

1120 **9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION**

1121

1122 Date of first authorisation: 12 February 2018

1123

1124

1125 **10. DATE OF REVISION OF THE TEXT**

1126

1127 (As per approval date of TH FDA)