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1 A randomised, open-label, parallel group phase 2 study of antisense oligonucleotide therapy in
2 acromegaly

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50 Key words: Acromegaly, antisense therapy, IGF-I, GH receptor

51

52 **Abstract**

53 Objective:

54 ATL1103 is a second-generation antisense oligomer targeting the human GH receptor. This phase 2
55 randomised, open-label, parallel-group study assessed the potential of ATL1103 as a treatment for
56 acromegaly.

57 Design:

58 26 patients with active acromegaly (IGF-I >130% upper limit of normal) were randomised to
59 subcutaneous ATL1103 200 mg either once- or twice-weekly for 13 weeks, and monitored for a further
60 8-week washout period.

61 Methods:

62 The primary efficacy measures were change in IGF-I at week 14, compared to baseline and between
63 cohorts. For secondary endpoints (IGFBP3, ALS, GH, GHBP), comparison was between baseline
64 and week 14. Safety was assessed by reported adverse events.

65 Results and Conclusions:

66 Baseline median IGF-I was 447 and 649 ng/mL in the once- and twice-weekly groups, respectively.
67 Compared to baseline, at week 14 twice-weekly ATL1103 resulted in a median fall in IGF-I of 27.8%
68 (p=0.0002). Between cohort comparison at week 14 demonstrated the median fall in IGF-I to be 25.8%
69 (p=0.0012) greater with twice-weekly dosing. In the twice-weekly cohort, IGF-I was still declining at
70 week 14, and at week 21 remained lower than at baseline by a median of 18.7% (p=0.0005).

71

72 Compared to baseline, by week 14 IGFBP3 and ALS had declined by a median of 8.9% (p=0.027) and
73 16.7% (p=0.017) with twice-weekly ATL1103; GH had increased by a median of 46% at week 14
74 (p=0.001). IGFBP3, ALS and GH did not change with weekly ATL1103. GHBP fell by a median of
75 23.6% and 48.8% in the once- and twice-weekly cohorts (p=0.027 and p=0.005), respectively.

76 ATL1103 was well tolerated, although 84.6% of patients experienced mild to moderate injection-site
77 reactions (ISR).

78 This study provides proof-of-concept that ATL1103 is able to significantly lower IGF-I in patients with
79 acromegaly.

80

81 Funding: Antisense Therapeutics Limited (Melbourne, Australia)

82 **Introduction**

83 Acromegaly is a rare, chronic, life-shortening disease caused by hypersecretion of growth hormone
84 (GH), virtually always due to a pituitary adenoma, that in turn results in elevated circulating levels of
85 insulin-like growth factor 1 (IGF-I).¹ Conventional therapy is directed at the pituitary gland and attempts
86 to reduce GH secretion by means of surgery, radiotherapy, or medical therapy in the form of
87 somatostatin analogues and dopamine agonists.² The GH receptor antagonist pegvisomant has
88 successfully exploited an alternative therapeutic approach, namely to block GH action rather than
89 secretion.³

90 Antisense oligonucleotides (ASOs) are single-stranded synthetic oligonucleotides that have been
91 developed as therapeutic agents. Translation of messenger RNA (mRNA), and hence protein synthesis,
92 is inhibited by sequence-specific ASOs which bind target pre-mRNA and/or mRNA.⁴ In the early
93 1990s, clinical trials with ASOs began, and in 1998 fomivirsen became the first oligonucleotide to be
94 approved by the U.S Food and Drug Administration (FDA) for the treatment of cytomegalovirus
95 retinitis.⁵ In 2013, the second-generation ASO inhibitor mipomersen was approved by the FDA for the
96 treatment of homozygous familial hypercholesterolaemia. Currently, there are more than 30 second-
97 generation ASOs, including ATL1103, in clinical development for a variety of neurological,
98 oncological, cardiovascular, and metabolic conditions. Excellent reviews of the technology are
99 available elsewhere.⁶

100

101 ATL1103 is a second-generation, antisense oligomer designed to inhibit translation of human growth
102 hormone receptor (GHR) mRNA (Figure 1). It comprises 20 nucleotides with a phosphorothioate
103 backbone and 2'-O-methoxyethyl modifications of the terminal five nucleotides at each end, which in
104 combination increase its plasma half-life and affinity for the mRNA. Post-hybridization RNaseH
105 degradation results in inhibition of GHR translation. In pre-clinical rodent and primate studies,
106 ATL1103 reduced GHR mRNA levels in the liver and serum IGF-I, with a terminal half-life of 2 to 4
107 weeks (add Tachas JoE 2006 189 147) Phase 1 studies in healthy male volunteers demonstrated a fall
108 in serum IGF-I and growth hormone binding protein (GHBP)
109 (<https://www.asx.com.au/asxpdf/20111207/pdf/4234016x2cj5xn.pdf>).

110

111 The objectives of this study were to evaluate the safety, tolerability, and efficacy of ATL1103 in patients
112 with acromegaly. Serum IGF-I was the primary measure of efficacy, with the other components of the
113 IGF ternary complex, namely IGF-binding protein 3 (IGFBP3) and acid labile subunit (ALS) being
114 additional measures of disease activity. Circulating GH and GHBP, the cleaved extracellular component
115 of the GHR, were monitored to provide insight into the physiological implications of an ASO targeting
116 the GHR. For the primary efficacy variable, the null hypothesis of no percentage change in fasting
117 IGF-I levels from baseline to week 14 was tested for each treatment regimen.

118

119 **Subjects and Methods**

120

121 Study Design

122

123 This was a phase 2 randomised, open-label, parallel-group study of the safety, tolerability,
124 pharmacokinetics, and efficacy of two subcutaneous dosing regimens of ATL1103 in patients with
125 acromegaly (Figure 2).

126

127 Exclusion/Inclusion Criteria

128

129 Inclusion criteria

130 Patients who:

- 131 1. provided written informed consent in accordance with local regulations
- 132 2. were 18 to 80 years of age inclusive
- 133 3. had acromegaly due to pituitary adenoma (micro- or macroadenoma) identified by magnetic
134 resonance imaging (MRI)
- 135 4. had serum IGF-I level at screening >1.3 times the upper limit of normal (ULN)

- 136 5. had documented serum GH nadir levels >1 ng/mL at all test time points within the 2 hours post
137 oral glucose load for an oral glucose tolerance test (OGTT) (this could be historical)
- 138 6. were acromegaly treatment naïve, or who had not taken other acromegaly medications for at
139 least the following periods of time prior to IGF-I and GH screening tests: bromocriptine: 6
140 weeks; carbergoline: 8 weeks; quinagolide: 8 weeks; octreotide (subcutaneous): 4 weeks;
141 pegvisomant: 8 weeks; octreotide LAR: 4 months; lanreotide (all presentations): 4 month
- 142 7. had a body mass index (BMI) ≥ 19 kg/m²
- 143 8. had adequate venous access to allow collection of multiple blood samples during the study
- 144 9. were female of non-child-bearing potential (i.e., either surgically sterilised or at least 1 year
145 post-menopausal), or, if of child-bearing potential, agreed to use two approved methods of
146 contraception for the duration of the study and for 3 months after administration of the last dose
147 of study drug; or were male and surgically sterilised or agreed to use an approved method of
148 contraception for the duration of the study and until 3 months after administration of the last
149 dose of the study medication
- 150 10. were willing and able to self-administer subcutaneous injections

151

152 (Inclusion criteria 5, 6, and 9 were amended in protocol amendments during the study. For inclusion
153 criterion 5, the requirement for GH after OGTT at screening was altered, as this could be historical.
154 Inclusion criterion 6 was amended to clarify that the washout periods detailed were minimum periods.
155 For inclusion criterion 9, contraceptive requirements were clarified.)

156

157 Exclusion criteria

158 Patients who:

- 159 1. had acromegaly due to reasons other than pituitary adenoma
- 160 2. had a pituitary adenoma that was less than 3 mm distance from the optic chiasm
- 161 3. had undergone pituitary surgery within the 3 months preceding the Screening visit
- 162 4. had received pituitary radiotherapy within the 1 year preceding the Baseline visit

- 163 5. had insulin-treated diabetes, or had commenced a new hypoglycaemic drug for diabetes within
164 the 2 months prior to Screening
- 165 6. had congestive heart failure, unstable angina, clinically significant cardiac arrhythmia, or a
166 history of acute myocardial infarction within the 3 months preceding the Baseline visit
- 167 7. had abnormal hepatic function at Screening defined any of the following parameters >2 x ULN:
168 aspartate aminotransferase (AST), alanine aminotransferase (ALT), gamma glutamyl
169 transferase (GGT), alkaline phosphatase (ALP), prothrombin time or total bilirubin
- 170 8. had hepatitis B, hepatitis C, or chronic liver disease
- 171 9. were pregnant or lactating
- 172 10. had known human immunodeficiency virus [(HIV) not tested specifically for this protocol], or
173 history of immunodeficiency that may have compromised their safety or affect results from this
174 study
- 175 11. had a history of alcohol or drug abuse in the 6 month period preceding the Baseline visit

176

177 Patients were recruited in 13 tertiary referral centres in Australia, France, Spain, and the United
178 Kingdom.

179

180 Appropriate ethical approval was obtained in every jurisdiction, and the study was registered as
181 EudraCT 201200314730 and ANZCTR 12611000854932. Patients gave written informed consent.

182

183 Procedures and Study Medication

184 Patients received either ATL1103 200 mg once or twice weekly 3 and 4 days apart for 13 weeks, with
185 every patient receiving three doses in the first week, administered every other day. Based on the tissue
186 half-life of >4 weeks, experience from primate studies and data from the phase 1 study, additional
187 'loading' doses were administered in the first week.

188

189 ATL1103 is formulated as a 'ready-to-inject' sterile solution at a concentration of 200 mg/mL, pH 7.4,
190 in 'Water for Injection'. Patients were taught to self-administer ATL1103 subcutaneously. After

191 completion of drug administration at the end of 13 weeks, patients were monitored, off all therapy for
192 acromegaly, for a further 8 weeks.

193 All patients underwent pituitary MRI scans at baseline and at week 13 (completion of the study drug),
194 which were independently reviewed by two 'blinded' expert pituitary neurosurgeons. Tumour diameter
195 changes of 2 mm or more in any one dimension or tumour volume changes of more than 20% were
196 considered significant.

197

198 An OGTT with measurement of plasma glucose and serum GH was undertaken at baseline (after any
199 drug washout) and again at the end of week 13. An adverse event (AE) assessment was undertaken at
200 each of the 11 study visits from baseline until study conclusion.

201

202 In addition to routine safety parameters, serum IGF-I, insulin-like growth factor binding protein 3
203 (IGFBP3), ALS, and GHBP were monitored. Ring size (fourth digit left hand) was measured using
204 standard European-sized jewelers' ring sets, and patients completed a signs and symptoms score (SSS,
205 maximum score 40) and the disease-generated 'quality of life' AcroQol. AcroQol comprises 22
206 questions divided into two main categories: physical and psychological function. The psychological
207 category is further subdivided into appearance and personal relationships. Each question is scored out
208 of 5, with a maximum score of 110 reflecting best possible quality-of-life. The result is then converted
209 to a percentage.⁷

210

211 Randomisation and Blinding

212 Permuted block randomisation (generated by a statistician and imported into the electronic case report
213 form) was used to assign patients to either open-label, once- or twice-weekly ATL1003. Once initial
214 data for a patient had been entered and the patient had fulfilled all inclusion criteria, a randomisation
215 number and treatment regimen were generated. Treatment allocation was not known to the operational
216 personnel until this randomisation was performed.

217 Blocks of size 4 were used with no stratification for the first 24 patients. The list included an

218 additional 24 randomisation numbers using a block size of 2 (total 48) to allow for overage.

219

220 Assays

221 IGF-I, GH, and IGFBP3

222 Serum IGF-I, GH, and IGFBP3 were measured centrally by IDS-iSYS (Immunodiagnostic Systems,
223 [IDS] Ltd., Boldon, England, UK) assays at the Endocrine Laboratory, Universität München (Munich,
224 Germany). Recombinant standards (98/574 for GH and 02/254 for IGF-I) yielded inter-assay
225 variability of 4.0–8.7% (IGF-I) and 1.1–3.4% (GH) and sensitivity of 8.8 ng/mL (IGF-I) and 0.04
226 ng/mL (GH).^{8,9} The limit of quantification for IGFBP3 was 50 ng/mL and the intra- and interassay
227 coefficients of variation (CVs) were 4.2% and 6.9%, respectively.¹⁰

228

229 ALS

230 Serum ALS levels were measured in duplicate by sandwich immunometric assay using monoclonal
231 antibodies directed against specific N- and C-terminal oligopeptides.¹¹ A serum pool of healthy male
232 volunteers was used for calibration and assigned 1000 U/litre. The assay range is 500 to 5000 U/litre,
233 and the intra- and interassay CVs were less than 9%. All samples from an individual subject were
234 analysed in one run.

235

236 GHBP

237 Serum GHBP levels were measured by a modification of the ligand immunofunctional assay with an
238 in-house monoclonal anti-GHBP antibody. Within-assay CVs were 9.4% at 115 pmol/L and 6.1% at
239 1550 pmol/L. At the same concentrations, between-assay CVs were 8.5% and 10.9%, respectively.
240 The lower limit of quantification was 69 pmol/L, and the linear range was 69–3500 pmol/L. All
241 samples from an individual subject were analysed in one run.¹²

242

243 Statistics

244 The study was powered for within-group comparison of serum IGF-I (primary efficacy variable).

245 Based on a published pegvisomant study,³ a clinically meaningful reduction in baseline IGF-I was
246 determined to be 27.5%, with a conservative estimate of standard deviation of 30%. To achieve a
247 level of significance of 0.05 with a two-sided test, it was determined that a minimum sample size of
248 12 patients per treatment was required to achieve a power of at least 80%.

249

250 The planned efficacy analyses were within the intention-to-treat group. For the primary efficacy
251 variable, the null hypothesis of no percentage change in fasting IGF-I levels from baseline to week 14
252 was tested for each treatment group with a (two-sided) one-sample *t* test and Wilcoxon signed rank
253 test. In addition, a pre-specified, though not powered for between-treatment-groups, comparison was
254 performed using a Wilcoxon Rank Sum test (mathematically equivalent to Mann-Whitney U test) for
255 serum IGF-I one week after the last dose of study drug (week 14).

256

257 Baseline to week 14 testing for both cohorts was undertaken (Wilcoxon signed rank test) for each of
258 the secondary endpoints (the other components of the IGF ternary complex, namely IGFBP3 and
259 ALS, plus circulating GH and GHBP, SSS, ring size, and AcroQoL) and presented as median plus
260 range.

261

262 A post-hoc regression analysis of the relationship between change in IGF-I and dose/kg/week is
263 reported with the associated 95% CI. Comparison of trapezoidal area-under-the-curve for GH during
264 the OGTTs is reported. Statistical significance is indicated by a p value <0.05.

265

266 Serum IGF-I data are expressed in mass units (ng/mL) and as a percentage of the upper limit of the
267 age-related reference range.

268

269 Data Safety Monitoring Board

270 An independent Data Safety Monitoring Board (DSMB) was established prior to recruitment start,
271 with an appropriate charter to direct decisions and monitor the trial safety results at intervals
272 throughout the study, and provided their recommendations as to whether the trial could continue as

273 planned or whether there were any concerns. The DSMB comprised four individuals with appropriate
274 experience in the areas of acromegaly, endocrine disorders, statistics, and the conduct of clinical
275 trials.

276

277 **Results**

278

279 Participants

280 Thirteen patients with active acromegaly (IGF-I >130% ULN at screening visit) (detailed in Table 1
281 and Figure 3) were recruited into each study arm.

282

283 Efficacy

284 IGF-I (Figures 4 and 5)

285

286 At baseline, the median serum IGF-I was 447 ng/ml (205–975)(2.5 x ULN) and 649 ng/ml (239–
287 831)(2.75 x ULN) ng/mL, in the once- and twice-weekly groups, respectively. Compared to baseline,
288 at week 14, ATL1103 at a dose of 200 mg twice weekly resulted in a median fall in serum IGF-I of
289 27.8% (range 4.4 to 49.8%, p=0.0002), while no change was seen with once-weekly dosing. At week
290 14, the median fall in IGF-I was 25.8% greater with twice-weekly compared to once-weekly dosing
291 (p=0.0012). In the twice-weekly cohort, IGF-I levels were still declining at week 14, and at the end of
292 the washout (week 21) remained lower than at baseline by a median of 18.7% (p=0.0005).

293

294 In both dosing regimens, one patient had an IGF-I within the age-related reference range at the pre-
295 defined endpoint of week 14. Normalisation of IGF-I at any time point was a pre-defined outcome
296 measure and was met by one additional patient in the twice-weekly regimen (week 13).

297

298 Combining the data from the two dosing regimens, regression analysis of the percentage change in
299 IGF-I levels versus dose/kg/week (median 2.88 (range 1.52 – 6.90) demonstrated an estimated slope
300 of regression of –8.27, indicative of a highly statistically significant (p=0.0001) association between

301 fall in IGF-I and the dose/kg/week (Figure 6).

302

303 IGFBP3

304 In the twice-weekly cohort, at week 14, there was a median fall in serum IGFBP3 of 8.9% (range -
305 29.2 to 12.9%, p=0.027) from baseline (median 7005 ng/ml, range 3396-9843). Once-weekly

306 ATL1103 did not result in a significant change in serum IGFBP3.

307

308 ALS

309 Compared to baseline, twice-weekly ATL1103 resulted in a median fall in ALS at week 14 of 16.7%
310 (range -20.9 to 34.9%, p=0.017) from baseline (median 1970 mU/ml, range 945-2463). Once-weekly

311 ATL1103 did not result in a significant change in serum ALS.

312

313 GH (Figure 7)

314 In the twice-weekly cohort, median trapezoidal AUC for GH during the OGTT had increased by 46%
315 (range -5.4 to 419%, p=0.001) at week 14 compared to baseline (471 ng.min/ml (79-867)). There was

316 no change in GH levels in the once-weekly cohort.

317

318 GHBP (Figure 8)

319 There was a significant decline in serum GHBP levels in both cohorts at week 14. Twice-weekly

320 ATL1103 resulted in a median decline of 48.8% (range -9.8 to 94.1%, p=0.005) in GHBP from
321 baseline (525 pmol/l (<69-6434)), while a median fall of 23.6% (range -61.4 to 59.4%, p=0.027) was

322 seen in the once-weekly cohort (1179 pmol/l (386-7637), p=0.027) and was maintained through to

323 week 21. In the twice-weekly cohort, the median fall in serum GHBP at week 21 was 40.4% (range -
324 94.1 to 6.1%, p=0.008) compared to baseline.

325

326 Ring Size

327 There was a statistically significant decrease in ring size circumference (mm) from baseline to week
328 13 for regimen 2, with a median decrease of -1.25 mm (range -12.6 to 3.8, p=0.039). Ring size was
329 unchanged with once-weekly dosing.

330

331 Signs and Symptoms Score

332 There was no marked difference in either regimen in median SSS at baseline (20 [1–36] vs 11 [7-30]).

333 The median percentage fall from baseline at week 14 was greater for twice-weekly dosing 37.5%

334 (range -185.7% to 91.7%) compared with 10.2% (range -33.3 to 83.3%) for once-weekly dosing,

335 although the changes were not statistically significant.

336

337 AcroQol

338 The median absolute improvement in the physical dimension and global scores between baseline and

339 week 14 in the once-weekly were 6.25 (range 0 to 31.3, p=0.002) and 3.4 (range -2.3 to 14.8,

340 p=0.0068) respectively, but these parameters did not change significantly with twice-weekly

341 ATL1103.

342 In contrast, in the twice weekly cohort, comparing baseline to week 14, the only significant finding

343 was an improvement in the median absolute change for the appearance subsection of psychological

344 dimension of 10.7 (range -17.9 to 25.0, p=0.035). There was no significant improvement in the once-

345 weekly cohort.

346

347 Safety (Table 2)

348 ATL1103 was well tolerated with mild to moderate injection site reactions (ISR) being the most

349 common, affecting 85% of patients in both cohorts, drug-related AE. Four serious AEs (SAEs) were

350 reported, of which three occurred in a single patient taking the once-daily regimen (acute bronchitis,

351 loss of consciousness while driving, and cholecystitis) and one in a patient taking the twice-daily

352 regimen (ear infection), but none were felt to be study drug-related, and both patients completed the

353 13 weeks of therapy. Two patients from one centre 'withdrew consent' at completion of dosing

354 (weeks 13 and 14) with study drug and so did not participate in the washout period through to week
355 21.

356

357 One patient in each regimen had low circulating platelet levels at a single time point (weeks 4 and 13,
358 86 and 132 x 10⁹/L, respectively; normal range 150 x 10⁹/L), but these resolved either spontaneously
359 or after treatment end (week 13). Two patients had elevated liver enzymes judged clinically
360 significant: one patient taking the once-daily regimen had γ -glutamyltransferase, AST, ALT, and ALP
361 values above the normal limit (>ULN); one patient taking the twice-daily regimen had AST and ALT
362 values >ULN. All effects on liver function were transient (Table 3).

363

364 The treatment-emergent AE (TEAE) profile was comparable for the two treatment groups
365 (Supplemental Table 2). Almost all patients experienced ISRs (mild and moderate), and 'mild'
366 lipohypertrophy, that subsequently, resolved was reported in two patients. There was a greater
367 incidence of headache in the once-weekly versus twice-weekly regimen (21 events and 5 events,
368 respectively), but the number of patients who experienced headache was comparable: 4 patients in the
369 once-daily regimen compared with 3 patients in the twice-daily regimen.

370

371 Radiologically significant tumour diameter changes (2 mm or more in any one dimension or tumour
372 volume changes of more than 20%) were reported in three patients. Tumour volume increased in two
373 patients (one in each dosing regimen, 5.7 x 7.3 x 19.1 v 6.8 x 9.9 x 19.5 mm and 8.1 x 5.8 x 14.8 v 8.1
374 x 7.2 x 16.2 mm) and reduced in one patient on twice-weekly dosing (6.2 x 10.4 x 4.9 v 2.6 x 5.7 x 4.1
375 mm). The changes were judged not to be clinically significant.

376

377 **Discussion**

378 The technology underpinning ASO therapy is rapidly advancing and has the potential to offer new
379 therapeutic options across a broad spectrum of diseases. Disordered protein production or function is
380 implicated in most pathological processes, and 'gene silencing or activating' by single-stranded

381 antisense oligonucleotides against target RNA sequences is an attractive concept that permits greater
382 specificity than can be achieved with small molecules or antibodies.¹³ The synthetic structural
383 modifications, such as the phosphorothioate backbone and 2'-O- methoxyethyl modifications, can be
384 readily applied to whole classes of ASOs with only the nucleotide sequence being indication specific.
385 Encouraging studies of ASOs are being reported against many targets, but this is the first report of the
386 use of an ASO in endocrinology.

387 The data presented provide the 'proof-of-concept' that in patients with acromegaly an ASO targeting
388 the GHR can lower serum IGF-I and raise the prospect of a new and entirely novel therapy for
389 acromegaly. Thirteen weeks of ATL1103 at a dose of 200 mg twice weekly lowered median serum
390 IGF-I by 27.8%, with two (15%) of 13 patients achieving an IGF-I within the reference range. Serum
391 IGF-I levels were still declining at week 14 and had not returned to baseline by the end of the 8-week
392 (week 21) washout period (Figure 4), suggesting that the duration of ATL1103 therapy was too short
393 to see maximum benefit and that prolonged treatment at the same doses may result in a further decline
394 in serum IGF-I. First-order drug kinetics indicate that approximately four to five elimination half-lives
395 are required to achieve steady-state plasma concentrations; since the tissue half-life of ATL1103 is
396 believed to be >4 weeks,¹⁴ this would suggest that between 16 to 20 weeks of treatment would be
397 required for nadir IGF-I levels to be achieved.

398

399 In conjunction with the data indicating a relationship between the dose per kilogram per week and the
400 fall in serum IGF-I, it seems probable that larger doses of ATL1103 administered for longer are likely
401 to result in a greater fall in IGF-I and offer the prospect of 'normalisation' of IGF-I in a greater
402 proportion of patients. Reassuringly, the decline in serum IGF-I with twice-weekly treatment was
403 paralleled by falls in the other elements of the IGF ternary complex, namely IGFBP3 and ALS.

404

405 Circulating GHBP is the product of cleavage of the extracellular component of the GH receptor.¹⁵ In
406 acromegaly there is a negative correlation between serum GHBP concentrations and IGF-I and GH
407 levels (Kratzsch Eur J Endocrinol 1995;132(3): 306e12), such that in active acromegaly GHBP

408 concentrations are decreased, and increase with conventional therapy. In contrast, the the reduction
409 in IGF-I caused by ATL1103 therapy is associated with a significant decline in serum GHBP
410 concentrations, which were still falling at week 14 and had not returned to baseline by the end of
411 washout at week 21. The fall in GHBP likely reflects the ATL1103-induced down-regulation of GHR
412 cell surface number, with a dose-response observed, as the fall was greater with the twice-weekly
413 compared to once-weekly regimen, 23.6% and 48.8%, respectively. The changes in GHBP emphasise
414 the difference in action of ATL1103 and future studies with increased doses of ATL1103 and larger
415 cohorts will permit exploration of the relationship between the change in serum IGF-I and circulating
416 GHBP concentrations.

417

418 As with pegvisomant, the reduction in serum IGF-I with twice-weekly ATL1103 was associated with
419 a 46% increase in serum GH levels assessed during an OGTT. Studies with pegvisomant have
420 demonstrated that the increased GH secretion is a consequence of negative feedback in response to the
421 fall in circulating IGF-I induced by blocking GH action.¹⁷

422 It is encouraging that an improvement in the soft-tissue manifestations of acromegaly, indicated by
423 the reduction in ring size, was seen with twice-weekly therapy. The short duration of therapy and
424 small cohort size means it should not be a surprise that the fall in IGF-I was not associated with an
425 improvement in SSS and only very modest improvements in quality of life as measured by AcroQoL.
426 Studies in larger numbers of patients treated for longer are required to demonstrate the impact of
427 ATL1103 on well-being and quality of life.

428

429 Almost all patients, approximately 85%, experienced ISRs, but otherwise ATL1103 was generally
430 well tolerated with no apparent drug-related SAEs. There were four SAEs, of which three occurred in
431 one patient, and all were judged to be unlikely to be drug related; both patients completed the 13
432 weeks of therapy (Table 2). ISRs are a recognised side effect of second-generation ASOs; were mild
433 to moderate in severity (predominantly a mixture of erythema, pain, and pruritus); and affected both
434 cohorts equally. No patient withdrew from this study because of ISRs. This is a similar incidence of

435 ISRs as reported in other studies: 90% of patients participating in a phase 2 study of mipomersen
436 experienced mild to moderate ISRs.¹⁸ The mechanism of oligonucleotide-induced ISR is yet to be
437 fully elucidated, but skin biopsies in 9 of 32 subjects participating in a phase 1 study of mipomersen
438 were consistent with leukocytoclastic vasculitis (e.g., infiltrating neutrophils, prominent nuclear dust,
439 lymphocytes, and eosinophils with local macrophage infiltration).¹⁹ The lessons learned from the
440 numerous other ASOs under clinical development will inform the management of ISRs in future
441 studies of ATL1103. Transient changes in platelet count and liver function were encountered but were
442 judged not to necessitate any change in therapy. Studies of greater length involving larger numbers of
443 patients are required before any conclusions can be drawn about the safety profile of ATL1103.

444

445 There were no clinically significant changes in pituitary tumour size, but the short duration of
446 treatment precludes meaningful conclusions about the long-term impact of ATL1103 on this
447 parameter. Both the patients in whom tumour expansion of >2 mm was documented had discontinued
448 somatostatin analogue therapy (one octreotide, one lanreotide) prior to commencing ATL1103, raising
449 the possibility of rebound expansion from somatostatin analogue-induced tumour shrinkage.
450 Reassuringly, the experience from more than 10 years' use of pegvisomant is that GHR-targeted
451 therapy does not induce growth of pituitary adenomas.²⁰

452

453 In summary, ATL1103 lowers IGF-I in acromegaly with biochemical changes consistent with down-
454 regulation of the GHR. As IGF-I was still declining at the end of the treatment period and with the
455 knowledge that the dose per kilogram could be increased, ATL1103 has the potential to achieve
456 disease control in a significant proportion of patients. Placebo-controlled studies of longer duration
457 and using higher doses are needed to better assess the full potential of this novel treatment.

458

459 **Declaration of interests**

460 PJT received research support from Antisense Therapeutics during the conduct of the study. JDCN-P
461 served as a consultant and steering committee member and received research support from Novartis.

462 PC received research support from Antisense Therapeutics, Novartis, Ipsen, Pfizer, and Italfarmaco.
463 TB received personal fees from Antisense Therapeutics and received research support and personal
464 fees from Pfizer, Ipsen, and Novartis. JA served as a speaker and on advisory boards for Pfizer, Ipsen,
465 and Novartis. GT and LA are employees of Antisense Therapeutics and own stock. MB received
466 research support from Antisense Therapeutics, research support and personal fees from Chiasma and
467 Novartis, and personal fees from ONO. All other authors have no relevant disclosures.

468

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471

472 **Author Contributions**

473 PJT contributed to the study design, identification of participating sites, and review of study data and
474 drafted the manuscript. JDCN-P participated in patient recruitment, contributed to drafting the
475 manuscript, and reviewed and approved the manuscript. JA, SJBA, AR, WD, PC, TB, SMW, CF, JA,
476 AIM, and DJT participated in patient recruitment and reviewed and approved the manuscript. GT and
477 LA were involved in protocol development and reviewed and approved the manuscript. DR was
478 responsible for the data analysis contained in the manuscript. MB contributed to the study design and
479 measurement of IGF-I, GH, IGFBP3, and ALS and reviewed and approved the manuscript.

480

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484

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557

558 **Tables**

559 **Table 1: Baseline Characteristics of Patients With Acromegaly**

	200 mg ATL1103 once weekly	200 mg ATL1103 twice weekly
	Median (range)	Median (range)
Number of patients	13	13
Age – years	49 (26–72)	49 (32–80)
Sex – M/F	5/8	6/7
Duration of disease* – years	9.0 (1–24)	3 (<1–20)
Weight – kg	90.6 (73.4–113.9)	83.2 (58–131.6)
Height – cm	169 (154–194)	163 (148–197)
Body mass index – kg/m ²	31.8 (26.0–39.6)	29.4 (21.4–45.3)
Size of adenoma at diagnosis – no. (%)		
Micro (<10 mm)	2 (18%)	2 (16.7%)
Macro (> or = 10 mm)	9 (81.8%)	10 (83.3%)
Missing	2	1
Hypopituitarism at study entry – no. (%)		
Surgery	13 (100%)	12 (92.3%)
Radiotherapy (all modalities)	6 (46.2%)	5 (38.5%)
Dopamine agonist therapy	3 (23%)	5 (38.5%)
Somatostatin analogue therapy	11 (84.6%)	12 (92.3%)

Pegvisomant therapy	7 (53.8%)	5 (38.5%)
Serum growth hormone - <i>ng/mL</i>	3.6 (0.4–60.6)	3.5 (1.5–9.4)
GH nadir (Screening OGTT) – <i>ng/mL</i>	2.5 (0.29–54.69)	2.4 (0.37–5.52)
Serum IGF-I – <i>ng/mL</i>	447 (205–975)	642 (239–831) [†]
Serum GHBP – <i>pM</i>	1179.0 (386–7637)	525.0 (<69–6434) [‡]
Serum IGFBP-3 – <i>ng/mL</i>	6589.0 (5162–9630)	7005.0 (3396–9843)
ALS – <i>mU/mL</i>	1669.0 (1395–2829)	1970.0 (945–2463)
Ring size circumference – <i>mm</i>	63.8 (57.5–81.4)	67.5 (53.7–78.9)
AcroQoL – <i>global score</i>	58 (18–100)	71 (56–90)
SSS – <i>calculated maximum score</i>	20.0 (1–36) [§]	11 (7–30)

560 GH, growth hormone; OGTT, IGF-I, insulin-like growth factor 1; GHBP, growth hormone binding
561 protein; ALS, acid labile subunit; IGFBP-3, insulin-like growth factor binding protein 3; SSS, signs
562 and symptoms score.

563 *Years from initial diagnosis to first day of study.

564 [†]Baseline IGF-I missing for one patient, screening IGF-I value used for calculations.

565 [‡]n=12, baseline GHBP missing for one patient.

566 [§]n=12, baseline SSS missing for one patient.

567

568

Table 2. Summary of Treatment-emergent Adverse Events (Safety Set)*

	200 mg once weekly (n=13)			200 mg twice weekly (n=13)			Total (N=26)		
	n	N	%	n	N	%	n	N	%
TEAEs	98	11	84.6	88	11	84.6	186	22	84.6
Drug-related TEAEs [†]	33	6	46.2	24	8	61.5	57	14	53.8
Serious TEAEs	3	1	7.7	1	1	7.7	4	2	7.7
Severe drug-related TEAEs	0	0		0	0		0	0	
Severe TEAEs	6	3	23.1	1	1	7.7	7	4	15.4
TEAEs leading to permanent discontinuation of study drug	0	0		0	0		0	0	
Withdrawals	1 [§]			1 [§]					
Patients with ISR	11		84.6	11		84.6	22		
Mild		9			6			15	
Moderate		2			5			7	
Severe		0			0			0	
Most frequent TEAEs with a >15% incidence									
Headache	21	4	30.8	5	3	23.1	26	7	26.9
Fatigue	3	2	15.4	3	2	15.4	6	4	15.4
Diarrhoea	3	2	15.4	2	2	15.4	5	4	15.4
Constipation	2	2	15.4	2	2	15.4	4	4	15.4
Dizziness	1	1	7.7	4	2	15.4	5	3	11.5
Hyperhidrosis	1	1	7.7	4	2	15.4	5	3	11.5
Rash	3	1	7.7	2	2	15.4	5	3	11.5
Abdominal pain upper	1	1	7.7	2	2	15.4	3	3	11.5
Nasopharyngitis	2	2	15.4	1	1	7.7	3	3	11.5
Urinary tract infection	3	2	15.4	0	0		3	2	7.7
Oropharyngeal pain	0	0		3	2	15.4	3	2	7.7
Abdominal distension	0	0		2	2	15.4	2	2	7.7
Abdominal pain	0	0		2	2	15.4	2	2	7.7

Tracheitis	2	2	15.4	0	0	2	2	7.7
Carpal tunnel syndrome	2	2	15.4	0	0	2	2	7.7
Haematuria	2	2	15.4	0	0	2	2	7.7
Lipohypertrophy	2	2	15.4	0	0	2	2	7.7

ISR, injection site reaction; N, number of patients; n, number of events; %, percentage of patients; TEAE, treatment-related adverse event.

*Excludes ISRs.

†Drug related is defined as relationship to study drug = possible, probable, or definite.

‡Withdrew consent after last drug dose.

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577 **Table 3. Summary of Abnormal Liver Function Tests.**

Analyte	Week	Result	Reference range
200 mg once weekly			
↑GGT, U/L	8	159 (repeats: 65, 32)	8–61
	21	102	
↑AST, U/L	8	111 (repeats: 70, 26)	6–40
↑ALT, U/L	8	181 (repeat: 135)	6–40
↑ALP, U/L	8	200	40–128
200 mg twice weekly			
↑AST, U/L	4	42	2–31
	8	43	
	4	69	
↑ALT, U/L	8	99	8–34
	13	52	
	1	28	
↑Total bilirubin (μmol/L)	2	22	0–21
	4	26	

578 GGT, gamma-glutamyl transferase; AST, aspartate aminotransferase; ALT, alanine aminotransferase;

579 ALP, alkaline phosphatase.

580

581 **Figures**

582

583 **Figure 1. Cartoon representation of ATL1103 and the mechanism of antisense inhibition.**

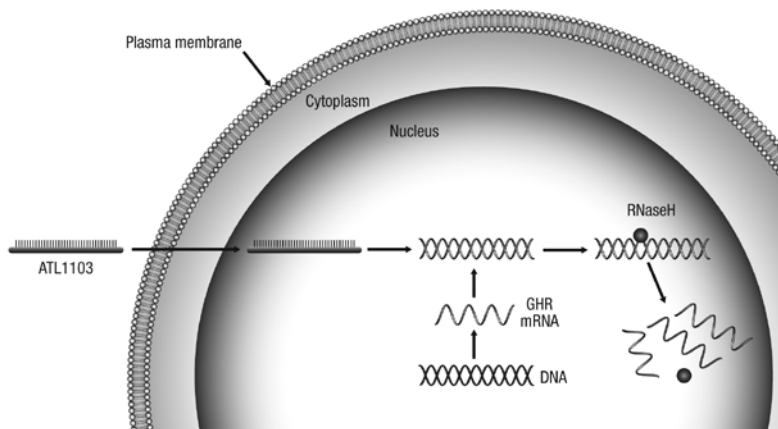
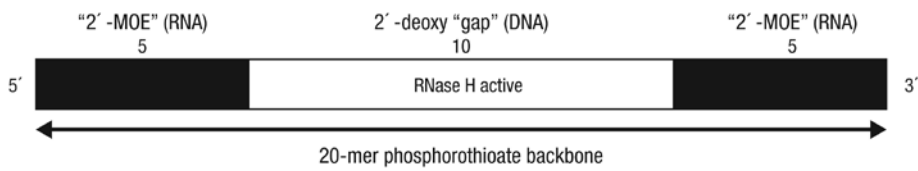
584 ATL1103 is a second-generation, 20-mer antisense oligonucleotide with a phosphorothioate backbone

585 and 2'-O-methoxyethyl modifications of the terminal five nucleotides at each end, which in

586 combination increase its plasma half-life and affinity for the mRNA. Post-hybridization RNaseH

587 degradation results in inhibition of GHR translation.

588



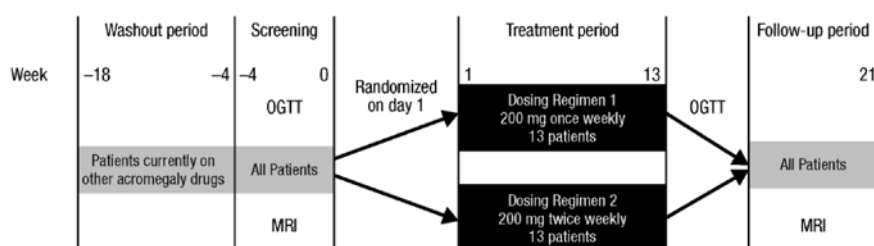
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590 GHR, growth hormone receptor.

591

592 **Figure 2. Schematic representation of study protocol.**

593 The protocol entailed appropriate washout from any ongoing acromegaly medical therapy after which
 594 serum IGF-I had to be at least >1.3 times age-related ULN. All patients underwent pituitary MRI
 595 scans at baseline and completion of the study drug. An OGTT was undertaken at baseline (after
 596 washout) and again at the end of week 13.



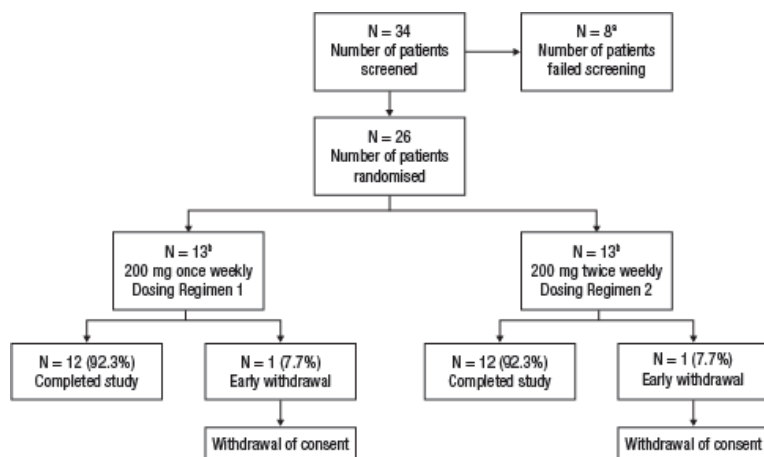
597
 598 IGF-I, insulin-like growth factor 1; ULN, upper limit of normal; MRI, magnetic resonance imaging;
 599 OGTT, oral glucose tolerance test.

600

601

602 **Figure 3. Patient disposition.**

603



604

605 ^aFive patients failed screening as IGF-I was <130% ULN.

606 ^bAlthough powered for 12 patients per arm, 13 were included per arm since a commitment had been

607 made to allow patients consented and 'passing' screening to receive study drug.

608 IGF-I, insulin-like growth factor 1; ULN, upper limit of normal.

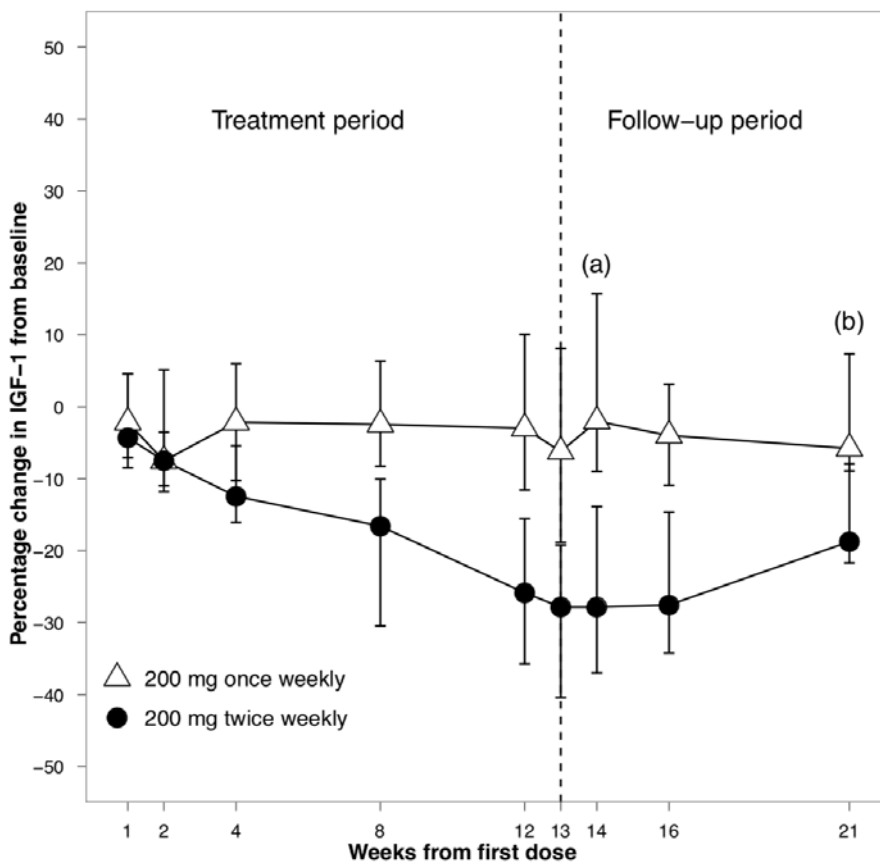
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611 **Figure 4. Median percentage change from baseline in serum concentrations of IGF-I in patients**
612 **with acromegaly.**

613 In the patients randomised to ATL1103 200 mg twice weekly, the median fall in serum IGF-I was
614 27.8% ($p=0.0002^a$) at the end of the treatment phase (week 14, 1 week after the last dose of study drug)
615 compared to baseline (week 0). Between cohort analysis at week 14 demonstrated the median fall in
616 serum IGF-I to be 25.8% ($p=0.0012$) with twice- compared to once-weekly dosing. In the twice-weekly
617 cohort, IGF-I was still declining at week 14, and at week 21 remained lower than at baseline by a median
618 of 18.7% ($p=0.0005^b$).

619



620

621

622 medians and interquartile ranges plotted

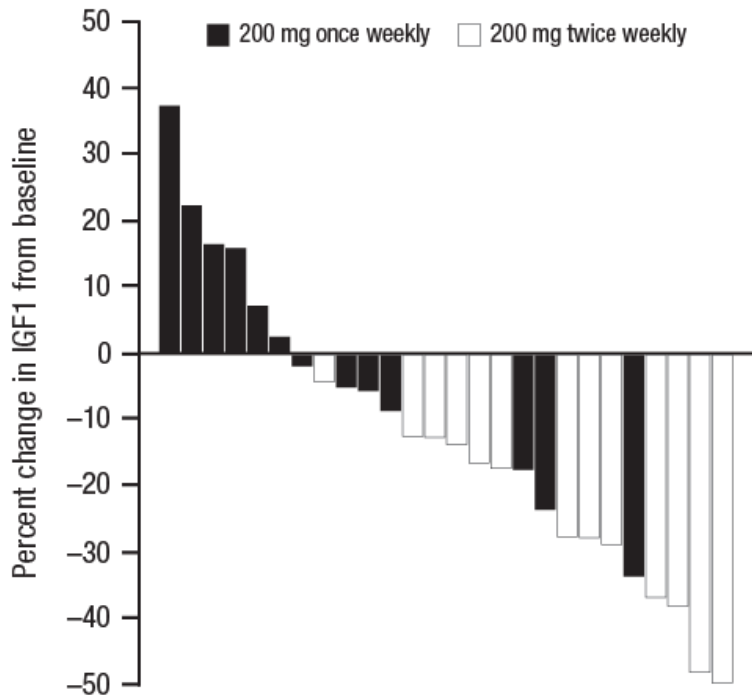
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627 **Figure 5. Percentage change in serum IGF-I levels from baseline to week 14 in 26 patients**
628 **treated with 200 mg ATL 1103 once or twice weekly.**
629



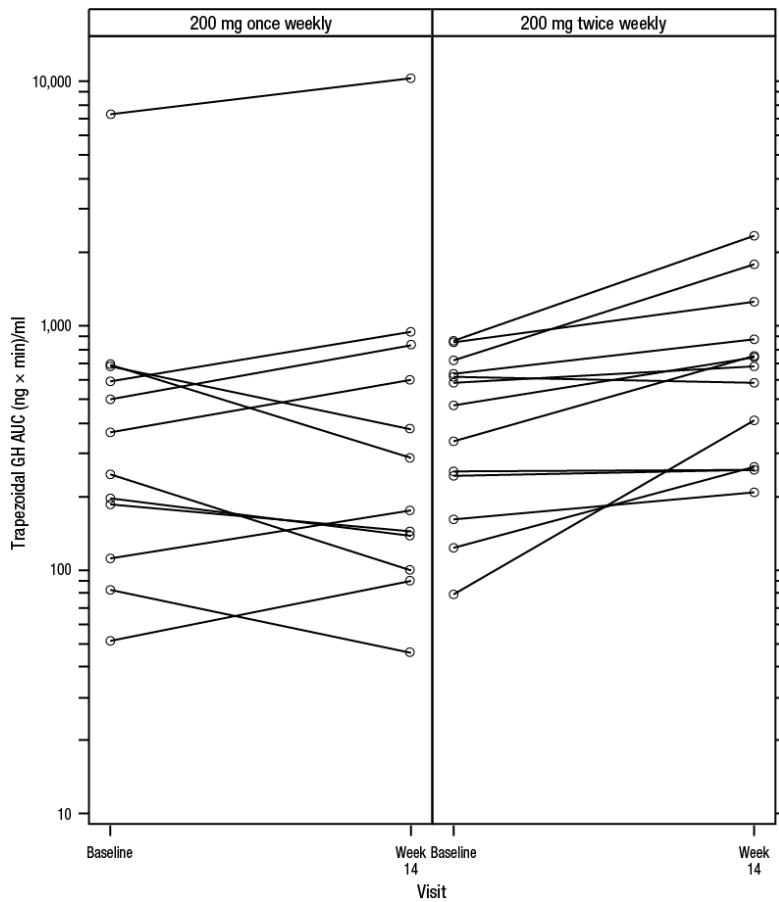
630
631 IGF-I, insulin-like growth factor .
632

639 **Figure 7. Trapezoidal AUC for GH during OGTTs at baseline and week 14.**

640 In the twice-weekly cohort, the median increase in AUC was 46% at week 14 compared to baseline
641 ($p=0.001$). There was no change in GH levels in the once-weekly cohort.

642

643

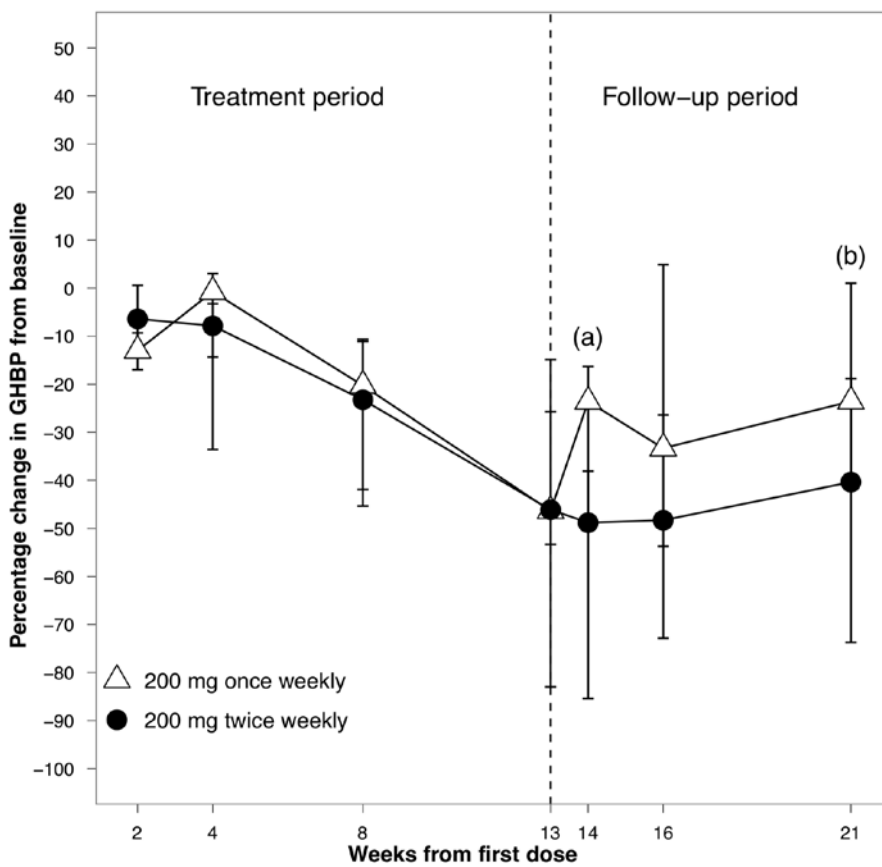


644

645 AUC, area under the curve; GH, growth hormone; OGTT, oral glucose tolerance test; CI, confidence
646 interval.

647

648 **Figure 8. Median percentage change from baseline in serum concentrations of GHBP in patients**
 649 **with acromegaly.**
 650 **Twice-weekly ATL1103 resulted in a median decline of 48.8% in GHBP ($p=0.005^a$) at week 14 (open**
 651 **symbols), while a median fall of 23.6% was seen in the once-weekly cohort ($p=0.027^a$). In the twice**
 652 **weekly cohort, at week 21 the median fall in GHBP, compared to baseline, was 40.4% ($p=0.008^b$).**



653
 654 **medians and interquartile ranges plotted**
 655