T2-weighted MRI signal predicts hormone and tumor responses to somatostatin analogs in acromegaly

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Abstract

GH-secreting pituitary adenomas can be hypo-, iso- or hyper-intense on T2-weighted MRI sequences. We conducted the current multicenter study in a large population of patients with acromegaly to analyze the relationship between T2-weighted signal intensity on diagnostic MRI and hormonal and tumoral responses to somatostatin analogs (SSA) as primary monotherapy. Acromegaly patients receiving primary SSA for at least

Key Words

- acromegaly
- somatostatin analogs
- MRI
- treatment response
- predictor

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3 months were included in the study. Hormonal, clinical and general MRI assessments were performed and assessed centrally. We included 120 patients with acromegaly. At diagnosis, 84, 17 and 19 tumors were T2-hypo-, iso- and hyper-intense, respectively. SSA treatment duration, cumulative and mean monthly doses were similar in the three groups. Patients with T2-hypo-intense adenomas had median SSA-induced decreases in GH and IGF-1 of 88% and 59% respectively, which were significantly greater than the decreases observed in the T2-iso- and hyper-intense groups ($P<0.001$). Tumor shrinkage on SSA was also significantly greater in the T2-hypo-intense group (38%) compared with the T2-iso- and hyper-intense groups (8% and 3%, respectively; $P<0.0001$). The response to SSA correlated with the calculated T2 intensity: the lower the T2-weighted intensity, the greater the decrease in random GH ($P<0.0001$, $r=0.22$), IGF-1 ($P<0.0001$, $r=0.14$) and adenoma volume ($P<0.0001$, $r=0.33$). The T2-weighted signal intensity of GH-secreting adenomas at diagnosis correlates with hormone reduction and tumor shrinkage in response to primary SSA treatment in acromegaly. This study supports its use as a generally available predictive tool at diagnosis that could help to guide subsequent treatment choices in acromegaly.

Introduction

Acromegaly is usually caused by a GH-secreting pituitary adenoma and has a prevalence of 8–12/100,000 people (Daly et al. 2006, Fernandez et al. 2010). Neurosurgical resection is the gold standard in acromegaly management (Katznelson et al. 2014), but remission after surgery alone is infrequent in macroadenomas, which constitute the majority of these tumors (Starke et al. 2013, Sarkar et al. 2014, Cuevas-Ramos et al. 2015). Consequently, medical treatment is often necessary and a variety of established and novel therapeutic options are available, including somatostatin analogues (SSA) (Melmed 2016). Responses to medical treatment are difficult to predict in advance of starting treatment and many current factors are based on histological characteristics of operated tumor tissue.

Pituitary MRI is routinely performed at diagnosis in patients with acromegaly. Apart from adenoma size and extension patterns, T2-weighted adenoma signal has recently been highlighted as a potential marker of disease behavior in acromegaly. In an international cohort of 297 patients with acromegaly, we showed that more than 50% of GH-secreting adenomas are T2-hypo-intense at diagnosis when compared to normal pituitary tissue (Potorac et al. 2015). These T2-hypo-intense adenomas are generally smaller, less frequently invade the cavernous sinus and are associated with higher levels of IGF-1. Interestingly, Puig-Domingo and coworkers previously noted that T2-hypo-intense post-operative adenoma residues ($n=62$) were more sensitive to subsequent SSA therapy (Puig-Domingo et al. 2010). Similarly, Heck and coworkers reported that T2-hypo-intense adenomas had a better hormonal response to pre-operatively administered SSA, suggesting that T2-hypo-intensity might be a useful pre-operative marker of SSA response (Heck et al. 2012, Heck et al. 2015). Recent guidelines for acromegaly management have also noted the potential utility of using T2 intensity to optimize patient management (Katznelson et al. 2014).

To assess the validity of this approach, we performed a multicenter, international study of patients with acromegaly and studied the relationship between the T2-weighted signal intensity on MRI and the hormonal and the tumor size responses to primary SSA therapy.

Patients and methods

This was an international study of acromegaly patients diagnosed from 2006 onwards at the 23 participating tertiary referral centers in 10 countries (Belgium, France, Germany, Italy, Russia, Spain, Netherlands, Turkey, Czech Republic and Algeria). All patients were diagnosed, managed and treated according to clinical criteria at their individual centers. Only patients with a visible GH-secreting pituitary adenoma of over 5 mm in diameter on diagnostic MRI and who had received primary SSA therapy for acromegaly were included. To be eligible, hormonal assessments and a follow-up MRI had to have been performed between 3 and 12 months after the beginning of primary treatment with SSA therapy.
The diagnosis of acromegaly was established based on an elevated IGF-1 level compared with the age- and sex-adjusted laboratory limits at each center; most patients (74%) also had evidence of an unsuppressed GH following an oral glucose tolerance test (OGTT). In patients who did not have an OGTT performed at diagnosis, IGF-1 level was more than two-fold higher than the upper limit of normal.

The study, its conduct and all of the analyses undertaken were performed under central approval for all sites by the Ethics Committee of the Centre Hospitalier Universitaire de Liège; this included anonymous data collection regarding the study participants.

Patient characteristics

The following data regarding each patient were recorded: date of birth, gender, age at diagnosis, type, dose and duration of SSA treatment. Each study participant’s data were blinded with a study code in order to maintain anonymity.

Hormonal assays

Values at diagnosis for random GH, GH nadir during OGTT, IGF-1 (absolute value and age and sex-adjusted upper limit of normal (ULN) of each laboratory) and prolactin were recorded. After 3–12 months of primary SSA treatment, a second hormonal evaluation (random GH, IGF-1 (absolute value and ULN), and prolactin) was performed. In order to eliminate variations between different assay kits and the different age- and sex-adjusted normal ranges, IGF-1 levels were expressed as a percentage of the ULN, further referred to as IGF-1%. Hormonal response to SSA treatment was evaluated as percent change from baseline in random GH and IGF-1% after SSA treatment. Co-secretion of prolactin was considered to be present when prolactin values exceeded 100 ng/mL at diagnosis, in accordance with the current literature (Karavitaki et al. 2006, Melmed et al. 2011).

Imaging criteria

The MRI examinations (at diagnosis and after 3–12 months of SSA treatment) of each patient were collected for central reading. This was performed by two experienced neuroradiologists and by consensus in cases of discordant readings. The T2-weighted signal on the diagnostic MRI was visually assessed as compared with normal pituitary tissue and when this was not visible, to the gray matter of the temporal lobe, as described previously (Potorac et al. 2015). Moreover, quantification of the T2-weighted signal by Region Of Interest (ROI) measurement of the adenoma, normal pituitary tissue and gray matter of the temporal lobe was performed and the calculated ROI-derived T2 intensity ratio of the adenoma vs gray matter was used for all further analyses.

We excluded adenomas that had a heterogeneous signal on diagnostic MRIs that made ROI measurement of T2-weighted signal intensity of the solid part of the adenoma unreliable. This allowed a uniform and unequivocal grading of the adenoma signal intensity and a consistent analysis based on this quantification. Very small adenomas with a maximum diameter <5 mm were also excluded as measurement of such small tumor sizes could have led to imprecise volumetric calculations.

For evaluation of adenoma size and tumor response to treatment, the tumor volume was calculated according to the formula: height × width × length × π/6 (Lundin & Pedersen 1992). Response to treatment was calculated as the percentage change in tumor volume between the two MRI examinations before and after SSA treatment.

Statistical analysis

Statistical analyses were performed using the R statistical package (R Core Team 2014). Graphs were generated using R and the Lattice graphical package (Sarkar 2008). Data were plotted and assessed for normal distribution. Since none of the variables showed a normal distribution, population spread was described using median and interquartile ranges (25th and 75th percentiles). Count variables were tested with the Chi-square test. Continuous variables were compared using the Mann–Whitney and Kruskal–Wallis tests. Graphs of univariate data are plotted as boxplots, with the limits of the box showing the 25th and 75th percentiles, the central line representing the median and the whiskers 1.5 times the interquartile range of the data.

Results

Patient characteristics

A total of 144 patients met the inclusion criteria of the study. Of these, 24 were excluded because the heterogeneous character of the adenoma on T2-weighted sequences made ROI analyses unreliable. The analysis was
therefore performed on 120 cases. Gender distribution was balanced (62 females (51.6%)). The median age at diagnosis of acromegaly was 51.0 years, with males being significantly younger than females (47.0 vs 59.0 years, respectively;  \( P = 0.0007 \)).

At diagnosis, the median random GH level was 9.9 ng/mL and the median nadir GH after glucose suppression was 5.5 ng/mL. The median IGF-1% at diagnosis was 314.9%; males had a trend toward higher IGF-1% values than females (335.3% vs 298.2%, respectively;  \( P = 0.06 \)). Prolactin co-secretion was found in eight patients.

Regarding SSA treatment, 45 patients received lanreotide Autogel (among these, one was switched to octreotide LAR), 69 patients received octreotide LAR (among them, two were switched to lanreotide Autogel) and three received pasireotide. For three patients, the SSA dose used was unknown. The duration of treatment between diagnostic and follow-up MRIs was similar between patients on lanreotide Autogel (5 months) and octreotide LAR (6 months). The median cumulative treatment doses were 540 mg for lanreotide Autogel and 150 mg for octreotide LAR.

Following SSA treatment, median random GH values were 2.3 ng/mL, and the median IGF-1% level was 146.8%. In total, 28 patients only (23%) had IGF-1 normalization after this short-term treatment period. Male patients had significantly higher IGF-1% levels on SSA treatment (197.4%) than females (120.1%;  \( P = 0.001 \)).

**Tumor characteristics on imaging**

There were 109 macroadenomas and 11 microadenomas. At the baseline MRI, the median adenoma volume was 1266 mm\(^3\) (Q1: 687 mm\(^3\), Q3: 2999 mm\(^3\)). The visual assessment of T2-weighted adenoma signal categorized 84 adenomas as hypo-intense, 17 as iso-intense and 19 as hyper-intense. Quantitative measurement of adenoma signal compared with normal pituitary tissue and gray matter confirmed the visual classification (Fig. 1). Using the ROI calculation, the adenomas could be clearly segregated into the three distinct groups with a clear demarcation among them that corresponded well with the visual classification. The ROI adenoma/ROI gray matter ratio for hypo-intense adenomas was 0.76 (Q1: 0.61, Q3: 0.86), while it was 1.05 (Q1: 1.02, Q3: 1.12) in iso-intense adenomas and 1.23 (Q1: 1.19, Q3: 1.38) in hyper-intense adenomas. Importantly, none of the adenomas that were hypo-intense on the visual classification had a ROI adenoma/ROI gray matter ratio \( \geq 1 \), and none of the hyper-intense adenomas on the visual classification had a ratio \( \leq 1 \).

Baseline hormonal parameters and adenoma volumes were significantly different between the groups. T2-hypo-intense adenomas had higher nadir GH values

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Baseline and post-treatment characteristics of the different T2-weighted signal groups of adenomas.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Hypo T2</td>
</tr>
<tr>
<td>Gender (M:F)</td>
<td>47:37</td>
</tr>
<tr>
<td>Age at diagnosis (years)</td>
<td>50.5</td>
</tr>
<tr>
<td>GH nadir (ng/mL) at diagnosis</td>
<td>10.6</td>
</tr>
<tr>
<td>IGF-1% at diagnosis</td>
<td>346.1%</td>
</tr>
<tr>
<td>Adenoma volume at diagnosis (mm(^3))</td>
<td>863</td>
</tr>
<tr>
<td>Random GH reduction (%)</td>
<td>88.0%</td>
</tr>
<tr>
<td>IGF-1% reduction (%)</td>
<td>59.1%</td>
</tr>
<tr>
<td>Volume reduction (%)</td>
<td>37.8%</td>
</tr>
</tbody>
</table>
and higher IGF-1% compared with T2-iso- and hyper-intense adenomas (Table 1). T2-hyper- and T2-iso-intense adenomas had significantly greater tumor volumes at diagnosis (3235 mm$^3$ and 2444 mm$^3$, respectively) as compared with T2-hypo-intense adenomas (863 mm$^3$; $P<0.0001$).

Response to treatment

Treatment with SSA was administered for a median duration of six months. There was no difference in median SSA treatment duration between the T2-hypo- (6 months), iso- (6 months) or hyper-intense (5 months) adenomas. Total cumulative and mean monthly doses of SSA were also similar between adenomas with T2-hypo-, iso- or hyper-intense signal intensities. No relationship was found between the decrease in GH or IGF-1% and the duration of treatment, total cumulative dose or mean monthly dose of either octreotide LAR or lanreotide Autogel. However, tumor shrinkage was found to be greater with longer treatment duration ($P=0.0003$, $r=0.11$) and with higher cumulative and mean monthly doses of octreotide LAR ($P=0.001$, $r=0.15$ and $P=0.0003$, $r=0.23$, respectively) and lanreotide Autogel ($P=0.02$, $r=0.13$ and $P=0.05$, $r=0.12$, respectively).

Hormonal response to SSA and T2-weighted signal

Using the visual assessment of T2-weighted signal, the decrease in random GH with SSA treatment was significantly greater in the T2-hypo-intense group (88.0%, Q1: 76.2%, Q3: 94.2%) as compared with T2-iso-intense (23.8%, Q1: 10.3%, Q3: 51.8%) and T2-hyper-intense (36.1%, Q1: 3.1%, Q3: 56.7%) groups ($P<0.0001$) (Fig. 2A). This decrease also correlated with the quantitative measurement of T2-weighted signal intensity (lower signal intensity adenomas had a greater GH decrease) ($P<0.0001$, $r=0.22$; Fig. 3A). The decrease in IGF-1% with

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**Figure 2**

Responses of random GH (A), IGF-1% (B) and volume (C) to SSA primary treatment depending on T2-weighted signal intensity.

**Figure 3**

Correlation between quantitative T2-weighted signal intensity of adenomas and their response to SSA in terms of GH (A), IGF-1% (B) and volume (C). GM, grey matter.
SSA treatment also was significantly greater in the visually assessed T2-hypo-intense (59.1%, Q1: 29.7%, Q3: 68.6%) as compared with T2-iso-intense (19.8%, Q1: 4.0%, Q3: 38.1%) and T2-hyper-intense (33.2%, Q1:13.0%, Q3: 47.9%) groups (P=0.0003) (Fig. 2B). As with GH, the quantitative measurement of T2-weighted signal intensity correlated with the decrease of IGF-1% (lower T2 intensity adenomas had a significantly greater IGF-1% decrease) (P<0.0001, r=0.14; Fig. 3B).

After SSA treatment, IGF-1 levels were normalized in 21/84 (25.0%) T2-hypo-intense, 7/19 (36.8%) hyper-intense and 0/17 iso-intense adenomas. Using a stricter criteria of IGF-1 control plus a random GH <1 ng/mL, disease control was achieved in 14/84 T2-hypo-intense (16.6%), no iso-intense and 3/19 hyper-intense (15.7%) adenomas (Fig. 4). No differences were seen in the statistical outcomes when the pasireotide-treated patients were excluded.

**Tumor shrinkage and T2-weighted signal**

There was significantly greater tumor shrinkage with SSA therapy in the T2-hypo-intense adenoma group (volume decrease: 37.8%, Q1: 24.7%, Q3: 49.3%) as compared with T2-iso-intense (8.0%, Q1: 4.1%, Q3: 13.8%) and T2-hyper-intense (3.0%, Q1: 1.0%, Q3: 13.3%) adenomas (P<0.0001; Figs 2C and 5). Quantitative measurement of adenoma signal intensity confirmed the visual findings and found a significant positive correlation between T2 intensity and volume reduction with lower intensity adenomas having a significantly greater volume reduction (P<0.0001, r=0.33; Fig. 3C). When considering the generally
accepted significant cut-off of >20% tumor shrinkage, 82.1% T2-hypo-intense adenomas, 15.8% T2-hyper-intense and 17.6% T2-iso-intense ones had achieved a significant volume reduction. Significantly more T2-hypo-intense adenomas, therefore, had a >20% tumor shrinkage compared with the T2-hyper- and iso-intense group (P<0.0001).

Discussion

In this multicenter study, we show that GH-secreting T2-hypo-intense pituitary adenomas at diagnosis have a significantly better response to primary SSA treatment not only in terms of hormonal but also anti-tumoral effects. T2-hypo-intense adenomas had the greatest hormonal response to SSA, with median decreases in random GH and IGF-1% of 88% and 59%, respectively. Furthermore, the shrinkage of T2-hypo-intense adenomas was of the order of 38%, whereas T2-iso- and hyper-intense adenomas had a median shrinkage <10%. These reductions in hormone levels and tumor size in T2-hypo-intense adenomas were greater than those seen with T2-iso-intense or T2-hyper-intense adenomas.

These results confirm and build upon previous studies on T2-weighted MRI signal intensity in acromegaly. As shown previously, T2-hypo-intensity is more frequent in smaller adenomas that are less invasive (Hagiwara et al. 2003, Heck et al. 2012, Potorac et al. 2015). Also, T2-hyper-intensity is more frequent in sparsely granulated pituitary adenomas (Hagiwara et al. 2003, Heck et al. 2012). As these characteristics are related to easier control of acromegaly, this has spurred interest in the relationship between T2 intensity and response to SSA therapy. Predictors of responses to therapy in acromegaly could be useful to match treatments to the needs and characteristics of the patient. As patients may undergo neurosurgery or medical therapy as a primary treatment, different predictors of remission and control are needed. Neurosurgical remission rates are related to lower preoperative GH levels, smaller tumor size, lack of invasion and the expertise of the surgeon (Ahmed et al. 1999, Nomikos et al. 2005, Starke et al. 2013). Recently, some novel genetic factors have been identified that predict difficult disease management in acromegaly. These newer factors, taken together with the standard pathological measures of tumor aggression such as granulation pattern, Ki-67 and gsp mutations can help to predict tumor responses to SSAs used post-operatively (Barlier et al. 1998, Bhayana et al. 2005, Fusco et al. 2008). For instance, mutations in the aryl hydrocarbon receptor interacting protein (AIP) gene lead to a significantly more aggressive phenotype in acromegaly including younger age at onset, larger tumor size, higher hormonal levels, and a significantly poorer response to SSA in terms of hormonal reduction and tumor shrinkage (Daly et al. 2010). Indeed, low immunohistochemical staining intensity for AIP in somatotrope tumors from acromegaly patients without AIP mutations is a predictor of a poor response to SSA (Jaffrain-Rea et al. 2009, Kasuki et al. 2012). X-linked acrogigantism (X-LAG) syndrome due to GPR101 duplication is also markedly SSA-resistant (Beckers et al. 2015, Rostomyan et al. 2015). Also, tumors with low levels of somatostatin receptor subtype 2 (SSTR2) tend to have a lower responsiveness to treatment with SSTR2-specific SSA like octreotide (Ferone et al. 2008, Taboada et al. 2008). Responses to pasireotide, a multi-somatostatin receptor specific ligand, also appear to be related to somatostatin receptor subtype 5 (SSTR5) expression (Iacovazzo et al. 2016). However, for predicting responses to primary pre-operative SSA therapy, non-invasive characteristics are required, which has driven the recent interest in MRI characteristics at diagnosis, particularly the T2 image intensity. It remains to be determined what relationship T2 intensity has to the underlying biological and pathophysiological nature of the tumor tissue and whether T2 signal intensity at baseline relates to immunohistochemical or genetic markers of tumor behavior in acromegaly.

In this study, the decreases in GH and IGF-1 correlated significantly with T2 signal intensity and the decreases were greatest among the T2-hypo-intense patients. These results are in keeping with those of Heck and coworkers, who showed median IGF-1 reductions of 51% for T2-hypo-intense, 36% for T2-iso-intense and 13% for T2-hyper-intense adenomas, respectively, and a higher IGF-1 control rate in T2-hypo-intense patients (Heck et al. 2012). In terms of control, however, in our study T2-hypo-intense adenomas had IGF-1 normalization in a quarter (21/84) of patients, whereas composite GH/IGF-1 control was achieved in 14/84 patients (16.6%). Normalization of IGF-1 and composite GH/IGF-1 was not achieved in any of the T2-iso-intense group, and in 7/19 and 3/19 patients, respectively, in the T2-hyper-intense group. It has been previously shown that lower baseline GH and IGF1 levels are associated with better hormonal responses to SSA treatment, expressed in terms of IGF1 normalization and achievement of a GH threshold concentration.
Here and previously we reported that T2-hypo-intense adenomas have higher baseline GH and IGF1 levels despite having smaller and less invasive tumors than non-T2 hypo-intense adenomas (Potorac et al. 2015). Here we show that these patients have greater percentage decreases from baseline in hormones and tumor size during primary SSA therapy. As they are starting from a higher hormonal baseline level, these patients frequently do not reach the threshold for strict control despite having greater percentage decreases from baseline IGF1. The control rate in T2 hypo-intense adenomas is, however, similar to that achieved in other ‘real world’ clinical treatment settings, in which less than 25% of patients achieve strict control with octreotide or lanreotide (Mercado et al. 2007, Espinosa-de-los-Monteros et al. 2015). For those T2-hypo-intense macroadenomas that appear difficult to fully resect with primary surgery, it may be that surgical debulking of such tumors would lead to a better hormonal control rate with post-operative SSA therapy (Petrossians et al. 2005, Jallad et al. 2007, Karavitaki et al. 2008).

Tumor shrinkage is an important benefit of SSA treatment in acromegaly that can alleviate clinical effects due to impingement on local structures around the pituitary tumor. Bevan and coworkers reported that around 50% of patients receiving primary SSA therapy have a significant volume reduction (Bevan 2005), whereas more recent studies report significant reductions after six months of treatment in 63% of patients (Mercado et al. 2007). The definition of significant volume reduction, however, varies from study to study and ‘significant’ shrinkage can be modest, of the order of 10–25%; for some years shrinkage of >20% has been considered to be clinically significant in trials of SSAs. We found that T2-hypo-intense tumors have significantly greater tumor shrinkage with SSA than T2-iso-intense or T2-hyper-intense adenomas. This shrinkage was nearly four times greater than in the other groups together and amounted to nearly 40%, which is a clinically meaningful change. When considering the >20% volume reduction criteria, more than 80% of T2-hypo-intense adenomas achieved this during SSA therapy as compared with less than 18% of either T2-hyper- and iso-intense adenomas. Previous studies of T2 intensity have shown variable results, with some indicating a correlation between T2 signal intensity at baseline and tumor shrinkage with SSA (Heck et al. 2015), whereas others showed no significant relationship (Puig-Domingo et al. 2010). This variability may have been due to the relatively small sample sizes analyzed for shrinkage in previous studies (n=34–49 patients), or due to methodological difficulties in measuring tumor shrinkage in previously operated tumor remnants (Puig-Domingo et al. 2010). The results of the current study indicate that T2-hypo-intensity is not only a predictor of hormonal responses but also of greater tumor shrinkage with SSA therapy in acromegaly.

A number of studies have assessed T2 signal intensity in acromegaly and its relationship to disease characteristics. While the results generally support the utility of T2 intensity as a predictor of therapeutic responses, some important methodological issues remain. In this and in our previous multicenter study we used a straightforward grading system for T2 intensity that is based primarily on the intensity of the adjacent normal anterior pituitary. We believe that this is the most relevant comparison, as it assesses the tumor tissue against unaffected normal tissue from the same gland. Only when such tissue cannot be visualized do we rely on the comparison with the temporal gray matter. In the vast majority of cases, however, normal pituitary tissue is readily visualized. This provides a single yardstick by which to judge hypo-, hyper- and iso-intensity. In this study we show that this simple visual classification correlates very closely with the precise but more time-consuming ROI analysis and allows for the three T2 intensity groups to be readily determined and with no inter-group overlap. This visual system, therefore, has the benefit of being reliable and easy to perform and provides an immediate assessment without further computerized analyses. Other classification systems rely on comparisons between the tumor characteristics and two non-pituitary signals, namely temporal white and gray matter. This raises problems when trying to compare the findings of different studies. For example, the response to SSA of T2-iso-intense adenomas differs between our study and the recent study of Heck and coworkers. For the reasons outlined above, we defined iso-intensity as a similar intensity to that of normal pituitary tissue or gray matter. In contrast, Heck and coworkers defined as iso-intense all adenomas that have T2 intensities intermediate between that of temporal white and gray (Heck et al. 2012). This leads to variability in the number of adenomas being assigned to the iso-intense group using the different classifications. Using our methodology, iso-intense adenomas accounted for only 14.2% of 120 acromegaly cases (70% were hypo-intense and 15.8% were hyper-intense), whereas all three groups...
were of similar proportion (28% hypo-intense, 36% iso-intense and 36% hyper-intense) in the Norwegian series. Moreover, depending on the technical characteristics of the MRI examination, contrast between gray and white matter can be adjusted, thereby modifying the range of intermediate intensities. In our study, T2 iso-intense adenomas are those adenomas with a very similar signal to that of gray matter and/or normal pituitary, as attested by the narrow corresponding range of ROI adenoma/ROI gray matter ratio (Fig. 1). Using our method, iso-intense adenomas represent a smaller group of tumors than using other methods, and their characteristics and behavior appear to be similar to the T2-hyper-intense group. This method to assess T2-weighted signal classification as compared with gray matter alone seems to be gaining popularity (Heck et al. 2015). From a practical point of view, we suggest that T2 iso-intense adenomas might in future be grouped along with hyper-intense tumors into a single non-hypo-intense category that has lower percentage decreases from baseline in hormonal levels and tumor volume. In support of this we found no statistically significant differences between the hormonal (in percentage decrease from baseline) and anti-tumoral response to SSA treatment of the T2 iso- and the hyper-intense groups. However, as the hormonal response of the iso- and hyper-intense groups did not separate completely from the T2 hypo-intense group in terms of IGF1 normalization, further work on this topic remains necessary.

A potential limitation of our study is the fact that it does not perfectly reflect the general population of acromegaly patients. Only adenomas treated pre-operatively with SSA were included. Therefore, the proportions of T2 hypo/iso/hyper-intense adenomas are not identical to those in large, unselected populations like in our previous study (Potorac et al. 2015). This point will require further studies to be performed on larger series of T2 iso- and hyper-intense adenomas, which could help to clarify the differences between T2 hypo-intense and non-hypo-intense tumors in acromegaly patients. Another potential issue is the difference in treatment timing and decision-making between centers. However, doses of SSA did not differ between adenomas with different types of T2-weighted signal, so treatments appear to be prescribed uniformly. In our view, having a heterogeneous international population of acromegaly patients from 23 centers may serve to reduce bias and indicates that the results could be applied to international acromegaly populations generally.

This large international study confirms that T2-weighted adenoma signal in acromegaly allows for the classification of adenomas into groups with different responses to SSA treatment. These differences are present in terms of both hormonal reduction and tumor shrinkage. The measured intensity of T2-weighted signal correlates with the magnitude of these responses; T2 hypo-intense adenomas constitute the largest group and have better hormonal and tumor shrinkage responses to SSA treatment. Visual assessment by comparison of adenoma signal to that of normal pituitary tissue or temporal lobe gray matter is reliable and easy to use in daily practice. These results from a diverse, international cohort support the utility of T2-weighted signal as a predictive factor of response to primary SSA treatment and may be a useful tool in future prospective studies of SSA in acromegaly.

Declaration of interest
The authors report no conflict of interest that could interfere with the performance, analysis or reporting of the study.

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References
Bhayana S, Booth GL, Asa SL, Kovacs K & Ezzat S 2005 The implication of somatotroph adenoma phenotype to somatostatin analog


R Core Team 2014 *R: A Language and Environment for Statistical Computing*. Vienna, Austria: R Core Team.


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