Multidimensional Geriatric Evaluation in Acromegaly: A Comparative Cross-Sectional Study

Irene Gagliardi  
University of Ferrara

Sabrina Chiloio  
Fondazione A Gemelli, IRCCS, Università Cattolica del Sacro Cuore

Maria Vallillo  
Fondazione A Gemelli, IRCCS, Università Cattolica del Sacro Cuore

Marta Bondanelli  
University of Ferrara

Stefano Volpato  
University of Ferrara

Antonella Giampietro  
Fondazione A Gemelli, IRCCS, Università Cattolica del Sacro Cuore

Antonio Bianchi  
Fondazione A Gemelli, IRCCS, Università Cattolica del Sacro Cuore

Laura De Marinis  
Fondazione A Gemelli, IRCCS, Università Cattolica del Sacro Cuore

Maria Chiara Zatelli  
University of Ferrara

Maria Rosaria Ambrosio (✉ mbmrs@unife.it)  
University of Ferrara

Research Article

Keywords: Acromegaly, Elderly, Cognition, Disability, Sarcopenia, QoL

DOI: https://doi.org/10.21203/rs.3.rs-514820/v1

License: ☒ This work is licensed under a Creative Commons Attribution 4.0 International License.  
Read Full License
Abstract

**Background:** Improvement in acromegaly management increased disease survival and prevalence. Evidence regarding acromegaly in older adults are sparse. We aim to explore acromegaly impact on aging process quality.

**Methods:** Multicenter case-control study conducted on 42 older adults (≥ 65 years) acromegaly patients (ACRO) compared to an age- and gender-matched control group (CTR). Each participant underwent a multidimensional geriatric evaluation.

**Results:** Mean age in both groups was 73 ± 6 years and female gender was most represented (69%). 13 ACRO were in remission and 29 had active disease controlled by medical therapy except for one patient. ACRO showed worse physical performance and mobility skills worsening with age as compared to CTR. ACRO performed poorly in functional status assessment, and age negatively correlated with instrumental and basic daily activities execution. Cognitive evaluation was significantly impaired in ACRO vs. CTR, worsening with age. No difference was found concerning nutritional and psychological status. Musculoskeletal and bone diseases were more frequent in ACRO than in CTR (52% vs. 12%; 64% vs. 10%; P< 0.05) and independently associated with geriatric outcomes in ACRO. ACRO reported a less satisfactory quality of life concerning physical activity and pain, general health, vitality, social activities.

**Conclusions:** Our study demonstrates increased frailty of older acromegaly patients as compared to non-acromegaly patients with a consequent negative impact on their quality of life. Therefore, it seems advisable to include physical, functional, cognitive, nutritional, and psychological status assessments in routine clinical practice. Further studies are needed to identify the most appropriate geriatric tools.

Background

Acromegaly is a rare disease caused by growth hormone (GH) hypersecretion mostly due to a pituitary adenoma, with consequent increase in insulin-like growth factor-1 (IGF-1) levels. Acromegaly presents with a progressive and chronic systemic involvement and comorbidities (cardiovascular, respiratory, metabolic, neoplastic, neurological, musculoskeletal) that are responsible for increased morbidity and mortality [1]. Literature reports a higher risk to develop cognitive impairment [2–6], psychopathologies [7–14], poor physical and functional performance [2, 15, 16] with negative effects on global quality of life (QoL). Improvement in diagnostic approaches, surgical techniques, medical therapies over the years has increased acromegaly survival and prevalence in all age groups [17–21]. Recent epidemiological data suggest a reduction in acromegaly mortality with death causes more similar to those of the general population of corresponding age. Therefore, the number of older patients with acromegaly, newly diagnosed or in follow-up, is expected to grow in the coming years. At the same time, the general population's life span increased [18–23]. We recently reviewed data from the literature to define acromegaly characteristics in the elderly but data regarding the impact of acromegaly disease on the natural aging course are still sparse [23]. Hatipoglu et al. have underlined the importance of a geriatric
multidimensional assessment also in older patients with acromegaly since the disease may enhance some aspects of the aging process (cognitive dysfunction, malnutrition risk, reduced physical performance, mood impairment) [2].

In this multicenter study we aim to define how acromegaly can impact on the quality of the aging process as compared to older general population. To this purpose, we performed a multidimensional evaluation of a cohort of Italian older acromegaly patients to assess their clinical, cognitive, functional, nutritional, and psychological status as compared to non-acromegaly older subjects.

**Methods**

This multicenter case-control study was conducted in two Italian endocrinological centers (Azienda Ospedaliero Universitaria of Ferrara and Policlinico A. Gemelli of Rome). Older acromegaly patients were recruited among those patients aged ≥ 65 years old. Acromegaly diagnosis was based on: 1) the presence of typical clinical features, 2) IGF-1 levels higher than age and sex adjusted reference ranges, 3) an abnormal GH response to an oral glucose tolerance test, 4) the presence of a pituitary adenoma. Study population included 42 cases (ACRO). The control group (CTR) included 42 age- and gender-matched non acromegaly subjects randomly selected among patients referred to an endocrinological and internist outpatient clinic (1:1 ratio). ACRO patients were divided in two groups according to the evidence of disease biochemical control at the last follow-up: patients in remission (R-ACRO) and patients with active disease (A-ACRO). R-ACRO patients presented normal IGF-1 levels for age and sex in the absence of medical treatment. A-ACRO patients, all receiving acromegaly medical treatment, were divided in two groups: patients with IGF-1 levels within the reference range (controlled A-ACRO) and patients with elevated IGF-1 (uncontrolled A-ACRO).

Demographic data, comorbidities, medications, and anthropometric measures were collected for each patient of both groups. For the ACRO group only, information about disease activity at last follow-up, previous and present acromegaly treatment, pituitary adenoma characteristics at diagnosis were gathered from clinical records. Cognitive function was assessed by using the age- and education-adjusted mini-mental state examination (MMSE) test [24]. Functional status was investigated with Lawton's instrumental activities of daily living (IADL) scale and with Barthel index for basic activities of daily living (BADL), that assess independence in daily living activities [25, 26]. Physical performance, mobility skills, and sarcopenia were tested by using the Timed Up and Go test (TUG), the Short Physical Performance Battery (SPPB) and the Handgrip test to both hands. Cut-offs established by The European Working Group on Sarcopenia in Older People (EWGSOP) were considered to interpret test results and to define sarcopenia presence and severity [27]. Nutritional status was determined by The Mini-Nutritional Assessment Scale-Short Form (MNA-SF), that screens for malnutrition risk [28]. Geriatric depression scale (GDS-15) by Yesavage [29] was used to screen the presence of depressive symptoms, evaluating the degree of depression. The SF-36 [30] test was used to investigate the self-reported measure of health-related quality of patient life, exploring the perception of eight life aspect domains. All tests and their
interpretation cut-offs are listed in Table 1. They were administered by two operators (one per center) equally trained in this study protocol.

This study is in accordance with the principles set out in the Declaration of Helsinki, has been specifically approved by the Local Ethics Committee (Comitato Etico Indipendente di Area Vasta Emilia Centro, CE-AVEC, at the Policlinico S.Orsola-Malpighi in Bologna) and authorized by the General Director of the Azienda Ospedaliero Universitaria in Ferrara (protocol number CE-AVEC 364/2018/Oss/AOUFe). All subjects read and signed the informed consent form before enrolling in the study.

**Statistical Analysis**

Categorical variables are presented as absolute values/percentages and Chi-square test was used to compare results. Continuous variables are described by mean values ± standard deviation (SD) and were compared by using the Student's T test in case of a normal distribution or by using the Mann–Whitney U test in case of non-normal distribution. Normality was tested with the Kolmogorov–Smirnov test. Statistical correlation between two continuous variables was evaluated by Pearson's correlation coefficient. P < 0.05 was considered as statistically significant. A multivariate analysis of variance (MANOVA) was performed to evaluate the involvement of comorbidites in the relationship between acromegaly and geriatric outcomes.
| Dimension assessed | Test | Cut-off and interpretation | References |
|--------------------|------|-----------------------------|------------|
| Cognitive function | MMSE | $\geq 24$: normal | 24 |
| | | $18-23$: mild cognitive impairment | |
| | | $11-17$: moderate cognitive impairment | |
| | | $<10$: severe cognitive impairment | |
| Functional status | IADL | $8 (5 \text{ in males})$: independence | 25 |
| | | $0$: total dependence | |
| | BADL | $100$: total independence | 26 |
| | | $91-99$: slight dependence | |
| | | $61-90$: moderate dependence | |
| | | $21-60$: severe dependence | |
| | | $0-20$: total dependence | |
| Physical performance | TUG | $<20 \text{ seconds}$: normal | 27 |
| Mobility skills | SPPB | $>8$: normal | |
| Sarcopenia | Handgrip test | $\geq 27 (\text{males})$: normal | |
| | | $\geq 16 (\text{females})$: normal | |
| Nutritional status | MNA-SF | $>11$: normal | 28 |
| | | $8-11$: risk of malnutrition | |
| | | $\leq 7$: malnutrition | |
| Depression | GDS-15 | $\leq 5$: normal | 29 |
| | | $6-10$: minor depression | |
| | | $>10$: major depression | |
| Quality of life | SF-36 | $100$: best own health perception | 30 |
| | | $0$: worse own health perception | |

**Results**

The mean age in both groups was $73 \pm 6$ years (range 65–91) and female gender was the most represented (69%). Daily medications number was significantly higher in ACRO as compared to CTR ($8 \pm 3$ vs. $3 \pm 2$), without significant differences regarding antidepressant/antipsychotic drugs. BMI was higher
in ACRO, but no significant differences were found concerning weight, waist and hip circumference as compared to CTR (Table 2).

| Clinical and anthropometric characteristics                  | ACRO     | CTR     |
|-------------------------------------------------------------|----------|---------|
| Total (n)                                                   | 42       | 42      |
| M/F                                                         | 13/29    | 13/29   |
| Mean age (years ± SD)                                       | 73 ± 6   | 73 ± 6  |
| Medications per day (n ± SD)                                | *8 ± 3   | 3 ± 2   |
| Antidepressants / antipsychotics / antiepileptic medications (n, %) | 11 (26%) | 5 (12%) |
| Weight (Kg ± SD)                                            | 79 ± 16  | 74 ± 10 |
| BMI (Kg/m² ± SD)                                            | *30 ± 6  | 28 ± 4  |
| • < 18.5 (n, %)                                             | 0 (0%)   | 0 (0%)  |
| • 18.5-24.99 (n, %)                                         | 6 (14%)  | 10 (23.8%)|
| • 25-29.99 (n, %)                                           | 17 (40%) | 22 (52.4%)|
| • 30-34.99 (n, %)                                           | 10 (24%) | 6 (14.3%)|
| • 35-39.99 (n, %)                                           | 5 (12%)  | 4 (9.5%) |
| • > 40 (n, %)                                               | 4 (10%)  | 0 (0%)  |
| Waist circumference (cm ± SD)                               | 137 ± 40 | 132 ± 39|
| Hip circumference (cm ± SD)                                 | 141 ± 37 | 139 ± 38|
| Waist/Hip circumference ratio                               | 0.97 ± 0.09 | 0.95 ± 0.09 |

* P < 0.05

ACRO clinical data at diagnosis and at last follow-up are summarized in Table 3.
Table 3
ACRO clinical data

| DIAGNOSIS                        |        |
|----------------------------------|--------|
| Mean age (years ± SD)            | 58 ± 10 |
| Age ≥ 65 (n, %)                  | 13 (31%)|
| Age ≤ 65 (n, %)                  | 29 (69%)|
| Microadenoma (n, %)              | 16 (38%)|
| Macroadenoma (n, %)              | 26 (62%)|
| Extra-sellar extension (n, %)    | 12 (29%)|
| LAST FOLLOW-UP                   |        |
| Remission (n, %)                 | 13 (31%)|
| Active controlled disease (n, %) | 28 (67%)|
| Active uncontrolled disease (n, %)| 1 (2%)  |
| Ongoing acromegaly therapy       |        |
| • None (n, %)                    | 13 (31%)|
| • SSA (n, %)                     | 15 (35.7%)|
| • PEG (n, %)                     | 2 (4.7%) |
| • DA (n, %)                      | 1 (2.4%) |
| • PEG + DA (n, %)                | 1 (2.4%) |
| • SSA + DA (n, %)                | 4 (9.5%) |
| • SSA + PEG (n, %)               | 5 (11.9%)|
| • SSA + PEG + DA (n, %)          | 1 (2.4%) |

SSA = somatostatin analogue; PEG = pegvisomant; DA = dopamine agonists

Among ACRO, 13 were in remission at the last evaluation (R-ACRO) and 29 presented active disease (A-ACRO), all controlled by medical therapy except for one patient with uncontrolled active disease. Concerning pituitary function, hypoadrenalism, hypothyroidism, hypogonadism and hyperprolactinemia were present in 21%, 50%, 17% and 10% of ACRO patients, respectively. Hyperprolactinemia was treated with dopamine agonists (DA). Replacement therapy was provided to all patients except for 3 hypogonadal male patients due to comorbidities. Cardio- and cerebro-vascular diseases, respiratory diseases, metabolic impairments, goiter, endocrine dysfunctions, gastrointestinal diseases, musculoskeletal complications, and bone diseases were significantly more frequent in ACRO as compared to CTR (Fig. 1).
Cognitive evaluation showed significantly lower MMSE scores in ACRO as compared to CTR (Table 4), without differences in cognitive impairment severity according to the test cut-offs. Scores negatively correlated with age in ACRO ($r = -0.25, P < 0.01$) and positively in CTR ($r = 0.37, P < 0.01$).

Functional assessment demonstrated significantly lower IADL and BADL scores in ACRO as compared to CTR, without differences in functional dependence levels. In ACRO both scores negatively correlated with age ($r = -0.29, P < 0.01$; $r = -0.41, P < 0.01$). On the contrary, in CTR age positively correlated with IADL ($r = 0.07, P < 0.01$) and negatively with BADL ($r = -0.29, P < 0.01$).

Regarding physical performance status and sarcopenia, ACRO obtained poorer results at TUG and SPPB tests as compared to CTR (Table 4). No difference was observed at the handgrip test and the strength developed by both hands did not differ between the two groups. In ACRO age negatively correlated with both SPPB test ($r = -0.37, P < 0.01$) and handgrip test (dominant hand $r = -0.29$ and non-dominant hand $r = -0.21, P < 0.01$). In addition, age positively correlated with time to complete the TUG test ($r = 0.17, P < 0.01$). In CTR only a negative correlation with the dominant handgrip test was found ($r = -0.18, P < 0.01$).

Nutritional status did not differ between groups on the basis of MNA-SF test scores. Scores become worse with age in ACRO but not in CTR ($r = -0.30$ vs. $0.06, P < 0.01$). Among ACRO, mean MNA-SF scores were significantly lower in R-ACRO than A-ACRO ($P < 0.05$), with a higher number of patients at risk of malnutrition as compared to A-ACRO ($69\%, P < 0.05$). Depression screening was negative in all subjects. GDS-15 scores did not show any correlation with age in both groups. All evaluations are summarized in Table 4.
Table 4
ACRO and CTR multidimensional evaluation

| Multidimensional evaluation | ACRO       | CTR       |
|-----------------------------|------------|-----------|
| **MMSE**                    |            |           |
| Score ± SD                  | *26 ± 5    | 28 ± 2    |
| Normal (n, %)               | 35 (83.3%) | 41 (98%)  |
| Mild cognitive impairment (n, %) | 5 (11.9%) | 1 (2%)    |
| Moderate cognitive impairment (n, %) | 1 (2.4%) | 0 (0%)    |
| Severe cognitive impairment (n, %) | 1 (2.4%) | 0 (0%)    |
| **IADL**                    |            |           |
| Score ± SD                  | *5 ± 2     | 7 ± 1     |
| Total independence (n, %)   | 20 (48%)   | 29 (69%)  |
| Total dependence (n, %)     | 0 (0%)     | 0 (0%)    |
| **BADL**                    |            |           |
| Score ± SD                  | *88 ± 23   | 99 ± 2    |
| Total independence (n, %)   | 27 (64.3%) | 37 (88.1%)|
| Slight dependence (n, %)    | 4 (9.5%)   | 2 (4.8%)  |
| Moderate dependence (n, %)  | 7 (16.6%)  | 3 (7.1%)  |
| Severe dependence (n, %)    | 2 (4.7%)   | 0 (0%)    |
| Total dependence (n, %)     | 2 (4.7%)   | 0 (0%)    |
| **TUG**                     |            |           |
| Seconds ± SD                | *16 ± 7    | 11 ± 5    |
| Sarcopenia (n, %)           | *14 (36%)  | 2 (5%)    |
| **SPPB**                    |            |           |
| Score ± SD                  | *7 ± 3     | 10 ± 2    |
| Sarcopenia (n, %)           | *28 (70%)  | 6 (15%)   |
| **HANDGRIP TEST**           |            |           |
| Dominant hand (Kg ± SD)     | 22 ± 10    | 24 ± 25   |
| Non dominant hand (Kg ± SD) | 17 ± 10    | 20 ± 8    |

*P < 0.05 ACRO vs. CTR
### Multidimensional evaluation

| ACRO       | CTR       |
|------------|-----------|
| Sarcopenia (n, %) | 14 (36%)   | 9 (21%)   |
| MNA-SF     |           |
| Score ± SD | 11 ± 2    | 12 ± 2    |
| Normal (n, %) | 22 (52%)  | 29 (69%)  |
| Malnutrition risk (n, %) | 18 (43%)  | 13 (31%)  |
| Malnutrition (n, %) | 2 (5%)     | 0 (0%)    |

### GDS-15

| ACRO       | CTR       |
|------------|-----------|
| Score ± SD | 5 ± 3     | 4 ± 3     |
| Normal (n, %) | 24 (57%)  | 30 (71%)  |
| Minor depression (n, %) | 16 (38%)  | 12 (29%)  |
| Major depression (n, %) | 2 (5%)     | 0 (0%)    |

*P < 0.05 ACRO vs. CTR

ACRO presented less satisfactory scores in 5 out of 8 SF-36 questionnaire domains as compared to CTR: physical activity, physical pain, general health, vitality, social activities (Fig. 2). Correlation with age was negative in all 8 fields in ACRO (P < 0.05). On the contrary, in CTR this correlation was significantly positive in the general health, vitality, and mental health fields. The multivariate analysis showed the significant involvement of specific comorbidities in negatively influencing geriatric outcome scores in ACRO (Table 5). No differences were observed comparing R-ACRO and A-ACRO in all items, except for MNA-SF scores.

### Table 5

Comorbidities significantly influencing geriatric outcome scores in ACRO by multivariate analysis

| Test | Comorbidities                                                                 |
|------|-------------------------------------------------------------------------------|
| IADL | Respiratory diseases, cardio and cerebrovascular disease, digestive diseases and bone diseases |
| BADL | Respiratory diseases, endocrine dysfunctions, digestive diseases and bone diseases |
| TUG  | Respiratory diseases, hypopituitarism, digestive diseases, bone diseases and musculoskeletal diseases |
| SPPB | Respiratory diseases, digestive diseases and bone diseases                  |

### Discussion
Our study demonstrates an increased frailty of older acromegaly patients as compared to non-acromegaly older patients, especially concerning cognitive function, functional status, and physical performance, with consequent negative impact on their quality of life.

In our study the ACRO group presented an average age of 58 ± 10 years at diagnosis. A third of them had been diagnosed after 65 years of age, possibly supporting the hypothesis that acromegaly has a mild clinical phenotype in the elderly patient [23, 31]. Two thirds were diagnosed before 65 years of age, therefore we cannot exclude an influence of disease duration on our results. Indeed, it is well consolidated that disease duration influences comorbidities development and worsening [32]. ACRO group was predominantly represented by females, in agreement with recent Italian series [20, 21, 33]. Pituitary adenomas were mostly macroadenomas, in contrast with literature data reporting that older patients generally carry smaller tumors as compared to younger patients [23]. This difference was further underlined by the finding that macroadenomas were more represented than microadenomas when considering only patients diagnosed with acromegaly after 65 years old. Therefore, our data indicate that age at diagnosis is not always predictive of tumor size. However, all macroadenomas were enclosed in the sellar space, in keeping with previous series [34–36] suggesting a milder behaviour of these tumors.

Our study shows a significant difference in BMI between ACRO and CTR, but waist/hip circumference ratio does not differ in the two groups, suggesting a similar fat tissue distribution. However, the available data do not allow to confirm this hypothesis and further studies are needed.

Most patients presented an active disease but controlled by medical treatment, mostly somatostatin analogs (SSA), in line with the guidelines [37]. Therefore, our ACRO group was homogeneous in terms of biochemical disease control, allowing us to hypothesize that the worse performance of ACRO vs. CTR was not due to aging nor to poor disease control, but possibly to the consequences of GH/IGF-1 excess before treatment. These data are in line with previous reports, showing that cognitive and psychological performance does not improve in acromegaly patients over time despite disease control [4, 9, 38, 39].

ACRO presented a more complex clinical picture than CTR, as indicated by the significantly higher number of medications per day and the more frequent comorbidities. Both acromegaly and aging are associated to comorbidities development, but our data suggest that acromegaly disease might be an additional factor negatively influencing older adults health. We observed a more important involvement of the musculoskeletal system in ACRO than in CTR. Indeed, arthropathy and osteoporosis are often described since the diagnosis of acromegaly and they usually persist also after remission [16, 40, 41]. Such evidence might partially explain our results derived from physical and functional evaluations. Indeed, ACRO obtained worse scores in the TUG and SPPB tests, describing worse physical performance and mobility skills. On the contrary, handgrip test results were similar in ACRO and CTR, in keeping with Füchtbauer et al. who did not find proximal muscle weakness in patients normalized after acromegaly remission [42]. We also found that bone and musculoskeletal diseases are independently associated with geriatric outcome scores in ACRO. Our results are in agreement with the study by da Silva Homem et al., who found a greater risk of falls and a worse performance concerning balance control, and peripheral
muscle function in 17 acromegalic older adults (age range: 63–73 years) compared with 20 paired controls [15]. Furthermore, we demonstrated that scores related to physical performance, mobility skills and sarcopenia worsen with age, suggesting a synergistic effect between aging and acromegaly, accounting for an increased frailty in older acromegaly patients. This hypothesis is further strengthened by the evidence that 3 out of 42 ACRO patients had been unable to complete the tests evaluating physical performance. Contrary to our findings, Hatipoglu et al. found no difference between the ACRO and CTR aging groups, by performing 2 out of 3 tests among those we employed. These differences may be due to the different number of evaluated patients; in addition, comorbidities were not fully characterized [2]. Furthermore, evidence regarding the relationship between acromegaly, sarcopenia and musculoskeletal status is still conflicting and very few data are available in the elderly, impairing the evaluation of the role of quantitative and qualitative muscle dysfunction [23].

Our study shows that functional status assessed by IADL and BADL is worse in ACRO vs. CTR, similarly to what found by Hatipoglu et al. [2]. Furthermore, in ACRO aging was negatively correlated with performance in instrumental and basic daily activities, possibly due to the worse musculoskeletal system condition and the presence of other comorbidities, as shown by the multivariate analysis. Our data support the hypothesis that musculoskeletal impairment is the primary source of physical pain and disability with negative impact on daily social activities. On the other hand, the employed evaluation was unable to discriminate the degree of dependence, in keeping with Fatti et al. [16] and suggesting that other tools are necessary to better identify those patients who need more assistance.

Cognitive dysfunction in acromegaly ranges from 2 to 33% of patients and is mostly represented by memory and attention impairments [43]. Neurocognitive alterations persist after surgical remission [4] suggesting that hormonal disease control is not sufficient to restore cognitive functions [38]. In keeping with these data, our experience shows that also in ACRO older patients cognitive performance is significantly impaired, despite disease control, worsening with age. Similarly to what found for functional status, MMSE was unable to capture the degree of cognitive impairment, probably due to the reduced size of our sample. Our data suggest that investigating cognitive function is appropriate, especially in older patients, considering MMSE scores as a preliminary evaluation.

Regarding nutritional and psychological evaluation, ACRO and CTR were similar in our settings. Hatipoglu et al., instead, described a worse nutritional status with a greater risk of malnutrition in acromegaly compared to the general older population, independently of GH/IGF-1 levels or disease activity [2]. These results suggest that MNA-SF and GDS-15 tests may not be sensitive enough to detect nutritional and psychological differences in controlled acromegaly versus normal aging subject. Further studies with larger cohorts and more sensitive tools are desirable.

Previous studies have widely documented the association between acromegaly and a poor QoL [6,39,44]. We found that QoL is worse in 5 out of 8 domains of the SF-36 questionnaire, confirming that acromegaly negatively impacts QoL also in the aging patient. AcroQoL, a disease-specific QoL questionnaire [45], is not applicable in our settings since it cannot measure QoL in CTR.
Finally, no relevant differences were found between A-ACRO and R-ACRO, but definitive conclusions cannot be derived due to the restricted number of the subsamples.

The multicenter design of our study allowed us to collect a significant number of older acromegaly patients, considering that acromegaly is a rare disease. Furthermore, two operators (one per center) equally trained in this study protocol have tested patients and controls in the two centers, making results more homogeneous. The multidimensional geriatric assessment of our study consists of simple tools readily available in routine clinical practice and extensively used in the geriatric general population. However, even if we derived significant conclusions from their applications, all tests did not perform equally in the acromegaly population. From our experience, the TUG and SPPB tests should be considered as useful tools to evaluate performance and sarcopenia status in acromegalic patients during routine clinical practice. They are easy to carry out and they are time and cost saving. However, body composition should also be explored besides patient performance, with specific tools (e.g. BIA, DEXA, MRI or CT).

**Conclusions**

Our multidimensional evaluation suggests that older acromegaly patients are much frailer than the general geriatric population, with a negative impact on their daily quality of life, supporting the indication to submit acromegaly patients to geriatric evaluation at an earlier age as compared to the general population. Our results suggest that disease management should take care also of such typical geriatric issues that could appear earlier and more importantly in acromegaly. Older acromegaly patients may indeed need assistance and support in their daily activities more often.

Therefore, it seems advisable to include physical, functional, cognitive, nutritional and psychological status assessments in routine clinical practice. However, other studies are needed to identify the most appropriate and efficient geriatric tools for acromegaly disease evaluation.

**Abbreviations**

| Abbreviation | Definition |
|--------------|------------|
| GH           | growth hormone; **IGF-1**: insulin-like growth factor-1; **ACRO**: patient group; **CTR**: control group; **MMSE**: mini-mental state examination test; **IADL**: Lawton's instrumental activities of daily living scale; **BADL**: Barthel index for basic activities of daily living; **TUG**: Timed Up and Go test; **SPPB**: Short Physical Performance Battery test; **MNA-SF**: Mini-Nutritional Assessment Scale-Short Form; **GDS-15**: Geriatric depression scale |

**Declarations**

**Ethics approval and Consent to participate**
This study was performed in line with the principles of the Declaration of Helsinki. Approval was granted by the Local Ethics Committee (Comitato Etico Indipendente di Area Vasta Emilia Centro, CE-AVEC, at the Policlinico S. Orsola-Malpighi in Bologna) and the study was authorized by the General Director of the Azienda Ospedaliero Universitaria in Ferrara (protocol number CE-AVEC 364/2018/Oss/AOUE). Informed consent was obtained from all individual participants included in the study.

**Consent for publication**

Patients signed informed consent for publication.

**Availability of data and materials**

The datasets generated and analyzed during the current study are available from the corresponding author on reasonable request.

**Competing interests**

The authors declare that they have no competing interests.

**Funding**

This study was supported by Funding from the University of Ferrara to MCZ and MRA (FAR2018).

**Author’s contribution**

Conceptualization: MRA; Methodology: IG. Investigation: IG, SC, MV; Formal analysis: IG; Writing - original draft preparation: MRA and IG; Writing - review and editing: MCZ, MB, SV, AG, AB, LDM; Supervision: MRA. All authors read and approved the final manuscript.

**Acknowledgments**

This work has been realized in collaboration with Laboratorio delle Tecnologie Avanzate (LTTA) at the University of Ferrara, Italy.

**References**

1. Colao A, Grasso LFS, Giustina A, Melmed S, Chanson P, Pereira AM, Pivonello. Acromegaly. Nat Rev Dis Primers. 2019;5:1-20. doi: 10.1038/s41572-019-0071-6.
2. Hatipoglu E, Yuruyen M, Keskin E, Yavuzer H, Niyazoglu M, Doventas A, Erdinciler DS, Beger T, Kadioglu P, Gundogdu. Acromegaly and aging: A comparative cross-sectional study. Growth Horm IGF Res. 2015;25: 47–52. doi: 10.1016/j.ghir.2014.12.003.

3. Leon-Carrion J, Martin-Rodriguez JF, Madrazo-Atutxa A, Soto-Moreno A, Venegas-Moreno E, Torres-Vela E, Benito-López P, Gálvez MA, Tinahones FJ, Leal-Cerro A. Evidence of cognitive and neurophysiological impairment in patients with untreated naive acromegaly. J Clin Endocrinol Metab. 2010;95:4367–79. doi: 10.1210/jc.2010-0394.

4. Martín-Rodríguez JF, Madrazo-Atutxa A, Venegas-Moreno E, Benito-López P, Gálvez MÁ, Cano DA, Tinahones FJ, Torres-Vela E, Soto-Moreno A, Leal-Cerro A. Neurocognitive Function in Acromegaly after Surgical Resection of GH-Secreting Adenoma versus Naïve Acromegaly. PLoS ONE. 2013;8:e60041. doi: 10.1371/journal.pone.0060041.

5. Sievers C, Sämann PG, Pster H, Dimopoulou C, Czisch M, Roemmler J, Schopohl J, Stalla GK, Zihl J. Cognitive function in acromegaly: description and brain volumetric correlates. Pituitary. 2012;15:350–7. doi: 10.1007/s11102-011-0326-z.

6. Szczęśniak D, Jawiarczyk-Przybyłowska A, Rymaszewska J. The Quality of Life and Psychological, Social and Cognitive Functioning of Patients with Acromegaly. Adv Clin Exp Med. 2015;24:167–72. doi: 10.17219/acem/38156.

7. Abed RT, Clark J, Elbadawy MH, Cliffe MJ. Psychiatric morbidity in acromegaly. Acta Psychiatr Scand. 1987;75:635–9. doi: 10.1111/j.1600-0447.1987.tb02848.x.

8. Richert S, Strauss A, Fahlbusch R, Oeckler R, von Werder K. Psychopathologic symptoms and personality traits in patients with florid acromegaly. Schweiz Arch Neurol Psychiatr Zurich Switz (1985). 1987;138:61–86.

9. Tiemensma J, Biermasz NR, van der Mast RC, Wassenaar MJE, Middelkoop HAM, Pereira AM, Romijn JA. Increased Psychopathology and Maladaptive Personality Traits, But Normal Cognitive Functioning, in Patients after Long-Term Cure of Acromegaly. J Clin Endocrinol Metab. 2010;95:E392–402. doi: 10.1210/jc.2010-1253.

10. Pereira AM, Tiemensma J, Romijn JA, Biermasz NR. Cognitive impairment and psychopathology in patients with pituitary diseases. Neth J Med. 2012;70:255-60

11. Alibas H, Uluc K, Kahraman Koytak P, Uygur MM, Tuncer N, Tanridag T, Yavuz DG. Evaluation of depressive mood and cognitive functions in patients with acromegaly under somatostatin analogue therapy. J Endocrinol Invest. 2017;40:1365–72. doi: 10.1007/s40618-017-0722-y.

12. Furman K, Ezzat S. Psychological features of acromegaly. Psychother Psychosom. 1998;67: 147–53. doi: 10.1159/000012275.

13. Sievers C, Ising M, Pfister H, Dimopoulou C, Schneider HJ, Roemmler J, Schopohl J, Stalla GK. Personality in patients with pituitary adenomas is characterized by increased anxiety-related traits: comparison of 70 acromegalic patients with patients with non-functioning pituitary adenomas and age- and gender-matched controls. Eur J Endocrinol. 2009;160:367–73. doi: 10.1530/EJE-08-0896.
14. Sievers C, Dimopoulou C, Pfister H, Lieb R, Steffin B, Roemmler J, Schopohl J, Mueller M, Schneider HJ, Isiing M, et al. Prevalence of mental disorders in acromegaly: a cross-sectional study in 81 acromegalic patients. Clin Endocrinol (Oxf). 2009;71:691–701. doi: 10.1111/j.1365-2265.2009.03555.x.

15. Homem TS, Guimarães FS, Soares MS, Kasuki L, Gadelha MR, Lopes AJ. Balance Control and Peripheral Muscle Function in Aging: A Comparison Between Individuals with Acromegaly and Healthy Subjects. J Aging Phys Act. 2017;25:218–27. doi:10.1123/japa.2016-0100.

16. Fatti LM, Cangiano B, Vitale G, Persani L, Mantovani G, Sala E, Arosio M, Maffei P, Dassie F, Mormando M, et al. Arthropathy in acromegaly: a questionnaire-based estimation of motor disability and its relation with quality of life and work productivity. Pituitary. 2019;22:552–60. doi: 10.1007/s11102-019-00966-8.

17. Ambrosio MR, Gagliardi I, Chiloiro S, Ferreira AG, Bondanelli M, Giampietro A, Bianchi A, De Marinis L, Fleseriu M, Zatelli MC. Acromegaly in the elderly patients. Endocrine. 2020;68:16–31. doi: 10.1007/s12020-020-02206-7.

18. Bolfi F, Neves AF, Boguszewski CL, Nunes-Nogueira VS. Mortality in acromegaly decreased in the last decade: a systematic review and meta-analysis. Eur J Endocrinol. 2018;179:9–71. doi: 10.1530/EJE-19-0509.

19. Lavrentaki A, Paluzzi A, Wass JAH, Karavitaki N. Epidemiology of acromegaly: review of population studies. Pituitary. 2017;20:4–9. doi: 10.1007/s11102-016-0754-x.

20. Burton T, Le Nestour E, Neary M, Ludlam WH. Incidence and prevalence of acromegaly in a large US health plan database. Pituitary. 2016;19:262–7. doi: 10.1007/s11102-015-0701-2.

21. Broder MS, Chang E, Cherepanov D, Neary MP, Ludlam WH. Incidence and prevalence of acromegaly in the united states: a claims-based analysis. Endocr Pract. 2016;22:1327–35. doi: 10.4158/EP161397.0R.

22. Arosio M, Reimondo G, Malchiodi E, Berchialla P, Borraccino A, De Marinis L, Pivonello R, Grottoli S, Losa M, Cannavò S, et al. Predictors of morbidity and mortality in acromegaly: an Italian survey. Eur J Endocrinol. 2012;167:189–98. doi: 10.1530/EJE-12-0084.

23. Caputo M, Ucciero A, Mele C, De Marchi L, Magnani C, Cena T, Marzullo P, Barone-Adesi F, Aimaretti G. Use of administrative health databases to estimate incidence and prevalence of acromegaly in Piedmont Region, Italy. J Endocrinol Invest. 2019;42:397–402. doi: 10.1007/s40618-018-0928-7.

24. Jallad RS, Bronstein MD. Acromegaly in the elderly patient. Arch Endocrinol Metab. 2019;63:638–45. doi: 10.20945/2359-3997000000194.

25. Folstein MF, Folstein SE, McHugh PR. “Mini-mental state”. A practical method for grading the cognitive state of patients for the clinician. J Psychiatr Res. 1975;12:189–98. doi: 10.1016/0022-3956(75)90026-6.

26. Shah S, Vanclay F, Cooper B. Improving the sensitivity of the Barthel Index for stroke rehabilitation. J Clin Epidemiol. 1989;42:703–9. doi: 10.1016/0895-4356(89)90065-6.
27. Lawton MP, Brody EM. Assessment of older people: self-maintaining and instrumental activities of daily living. The Gerontologist. 1969;9:179–86.

28. Cruz-Jentoft AJ, Bahat G, Bauer J, Boirie Y, Bruyère O, Cederholm T, Cooper C, Landi F, Rolland Y, Sayer AA, et al. Sarcopenia: revised European consensus on definition and diagnosis. Age Ageing. 2019;48:16–31. doi: 10.1093/ageing/afz046.

29. Kaiser MJ, Bauer JM, Ramsch C, Uter W, Guigoz Y, Cederholm T, Thomas DR, Anthony P, Charlton KE, Maggio M, et al. Validation of the Mini Nutritional Assessment short-form (MNA-SF): a practical tool for identification of nutritional status. J Nutr Health Aging. 2009;13:782–8. doi: 10.1007/s12603-009-0214-7.

30. Martínez de la Iglesia J, Onís Vilches MC, Dueñas Herrero R, Aguado Taberné C, Albert Colomer C, Arias Blanco MC. Abbreviating the brief. Approach to ultra-short versions of the Yesavage questionnaire for the diagnosis of depression. Aten Primaria. 2005; 35:14–21. doi: 10.1157/13071040.

31. Apolone G, Mosconi P. The Italian SF-36 Health Survey: translation, validation and norming. J Clin Epidemiol. 1998;51:1025–36. doi: 10.1016/s0895-4356(98)00094-8.

32. Gatto F, Trifirò G, Lapi F, Cocchiara F, Campana C, Dell’Aquila C, Ferrajolo C, Arvigo M, Cricelli C, Giusti M, et al. Epidemiology of acromegaly in Italy: analysis from a large longitudinal primary care database. Endocrine. 2018;61:533–41. doi: 10.1007/s12020-018-1630-4.

33. Melmed S, Bronstein MD, Chanson P, Klibanski A, Casanueva FF, Wass JAH, Strasburger CJ, Luger A, Clemmons DR, Giustina A. A Consensus Statement on acromegaly therapeutic outcomes. Nat Rev Endocrinol. 2018;14:552–61. doi: 10.1038/s41574-018-0058-5.

34. Prencipe N, Scarati M, Manetta T, Berton AM, Parisi S, Bona C, Parasiliti-Caprino M, Ditto MC, Gasco V, Fusaro E, et al. Acromegaly and joint pain: is there something more? A cross-sectional study to evaluate rheumatic disorders in growth hormone secreting tumor patients. J Endocrinol Invest. 2020;43:1661-1667. doi: 10.1007/s40618-020-01268-8.

35. Muzziotti G, Maffezzoni F, Frara S, Giustina A. Acromegalic osteopathy. Pituitary. 2017;20:63–9. doi: 10.1007/s11102-016-0758-6.

36. Füchtbauer L, Olsson DS, Bengtsson B-Å, Norman L-L, Sunnerhagen KS, Johannsson G. Muscle strength in patients with acromegaly at diagnosis and during long-term follow-up. Eur J Endocrinol. 2017;177:217–26. doi: 10.1530/EJE-17-0120.

37. Brummelman P, Koerts J, Dullaart RPF, van den Berg G, Tucha O, Wolffentuttel BHR, van Beek AP. Effects of previous growth hormone excess and current medical treatment for acromegaly on cognition. Eur J Clin Invest. 2012;42:1317–24. doi: 10.1111/j.1365-2362.2012.02721.x.

38. Yedinak CG, Fleseriu M. Self-perception of cognitive function among patients with active acromegaly, controlled acromegaly, and non-functional pituitary adenoma: a pilot study. Endocrine. 2014;46:585–93. doi: 10.1007/s12020-013-0106-9.

39. Crespo I, Valassi E, Santos A, Webb SM. Health-related quality of life in pituitary diseases. Endocrinol Metab Clin North Am. 2015;44:161–70. doi: 10.1016/j.ecl.2014.10.013.
40. Webb SM, Badia X, Surinach NL. Validity and clinical applicability of the acromegaly quality of life questionnaire, AcroQoL: a 6-month prospective study. Eur J Endocrinol. 2006;155:269–77. doi: 10.1530/eje.1.02214.