Effect of Biologic Treatments on Growth in Children with Juvenile Idiopathic Arthritis

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ABSTRACT. Objective. Growth retardation is a frequent complication of severe juvenile idiopathic arthritis (JIA). Biologic treatments may improve growth velocity by controlling systemic inflammation and reducing corticosteroids. Our goals were to compare growth velocity before and after the onset of biologic therapy and to determine whether the JIA subtype, the use of steroids, the requirement of one or several biologic agents, or the disease activity influenced growth velocity.

Methods. We retrospectively analyzed the growth of children with JIA who never received growth hormone treatment, who started biologic treatment before puberty, and who were followed for at least 6 months afterward.

Results. We included 100 children (33 boys). Median patient age was 7.1 years (range: 1.6–15.7) at the onset of biologic treatment and 11.0 years (range: 2.3–19.5) at the latest followup. Forty-six patients had received corticosteroid and 34 had received more than 1 biologic agent. Patient median height expressed as SD score (SDS) was 0.31 (range: –2.47 to 5.46) at disease onset, –0.24 (–3.63 to 2.90) at biologic therapy onset (p < 0.0001), and –0.15 (–4.95 to 3.52) at the latest followup (p = 0.171 compared to biologic treatment onset). Patients who required several biologics and systemic patients had a significantly lower growth velocity after the onset of biologic treatment. At the latest followup, 18% of our study group had low growth velocities and 19% were below –2SD or shorter than genetically programmed.

Conclusion. In a subset of patients, particularly systemic JIA patients and patients who required more than 1 biologic, biologic therapy may be insufficient to restore normal growth velocity.

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Key Indexing Terms:

JUVENILE ARTHRITIS SYSTEMIC JUVENILE ARTHRITIS CYTOKINE BIOLOGIC TREATMENT GROWTH

Juvenile idiopathic arthritis (JIA) is a heterogeneous group of diseases characterized by arthritis of unknown etiology starting before the age of 16 years and lasting for more than 6 weeks1. The prevalence of JIA has been estimated to be between 16 and 150 per 100,000 children2. JIA may be responsible for pain, joint destruction, functional impairment, and in some patients, short stature in adulthood3,4,5. According to the most recent classification (revised in 2001), there are 7 JIA subtypes: systemic-onset JIA (SJIA), characterized by the association of systemic features that include a spiking fever and arthritis; rheumatoid factor (RF)-positive polyarthritis; RF-negative polyarthritis; oligoarthritis, characterized by arthritis affecting < 5 joints during the first 6 months of the disease; enthesitis-related arthritis; psoriatic arthritis; and undifferentiated arthritis6.

In the most severe cases, and especially in children with SJIA, the combination of chronic inflammation and corticosteroid treatment can cause severe growth retardation. Conventional disease-modifying antirheumatic agents (DMARD), such as methotrexate (MTX), have been reported to improve JIA patient growth but their effect is
insufficient in many cases. Growth hormone (GH) therapy has been shown to improve and often normalize growth velocity in JIA patients who required long-lasting corticosteroid treatment. However, GH therapy remains experimental for this indication. Biologic treatments have been used in patients with JIA for more than 12 years. In addition to their powerful effects against joint inflammation and the possible prevention of joint erosion, they may suppress systemic inflammation and may have a corticosteroid-sparing effect in many cases. Some recent reports indicated that anti-tumor necrosis factor-α (anti-TNF) treatments were associated with growth velocity improvement in patients with JIA. In these studies, however, only a small proportion of the patients had SJIA, which is associated with severe systemic inflammation and high corticosteroid requirements, and hence carries the highest risk of growth failure. In addition, patient target height was not documented.

In our study, we assessed height and changes in the height SD scores at disease onset, at the onset of biologic therapy, and at the last follow-up visit in a cohort of JIA children. All patients were prepubertal or in early puberty when biologic treatment began and were followed for at least 6 months afterward. We conducted this study in a referral pediatric rheumatology center where hundreds of patients with JIA are followed, including a high proportion of patients with SJIA. By taking into account the target height of these patients, our aim was to better assess the effect of biologics on growth in patients with JIA and to analyze the proportion of patients in whom persistent growth impairment might justify GH treatment. We also sought to determine whether the height SD at the latest follow-up was influenced by JIA subtype, by comparing patients with SJIA to other patients with JIA, by the presence of corticosteroid treatment, by the requirement of > 1 biologic during the follow-up, and by achieving inactive disease at the latest follow-up.

**MATERIALS AND METHODS**

**Inclusion criteria.** Patients were required to fulfill the International League of Associations for Rheumatology (ILAR) criteria for JIA, to be under biologic therapy for at least 6 months so that it would be possible to assess changes in height SD score (SDS), and to be prepubertal or early pubertal at biologic treatment onset. We included all of the patients who met these criteria and were cared for at the French National Reference Center for Juvenile Arthritis at the Necker-Enfants Malades Hospital, Paris, France, between May and October 2010.

As documented in the patient notes, the parents and patients (when old enough) were informed and agreed that patient data could be recorded in a database built for patients with rare diseases, the CEMARA platform. This included an agreement from the French “Commission Nationale Informatique et Libertés”. In accordance with the French rules, no written informed consent was requested for this retrospective study.

**Exclusion criteria.** Patients who were postpubertal (Tanner staging > 3) at biologic therapy onset and patients with a joint deformity that would not allow proper height assessment were not eligible for the study. Patients with an associated medical condition that could modify their growth and patients who were receiving or had received GH therapy were excluded. Patients were also excluded if it was not possible to retrospectively collect enough data to properly assess their growth.

**Data collection.** Patient notes were reviewed in detail by one of the authors (FU) who, along with a reference pediatric rheumatologist (PQ, BBM, RM, or CW), also saw the patients and their parents on one or several occurrences during the 6-month inclusion period.

The following data were collected: height and weight at birth; height and weight at each outpatient clinic; pubertal stage at biologic treatment onset and at the last follow-up, according to Tanner classification; mother’s and father’s heights; patient JIA subset, according to the ILAR classification; presence/absence of uveitis; age at disease onset; and any other medical problems and their treatments. Age at disease onset was defined by patient’s age at the first clinical symptom (joint pain or limitation, swollen joint, uveitis, or systemic symptoms). According to Wallace criteria, we consider active disease in the form of 1 or more of the following: persistent active arthritis; disease activity that was determined by the physician to be above 10/100 on a visual analog scale; clinical systemic features, such as fever or pericarditis; active uveitis; and/or an elevated erythrocyte sedimentation rate or serum level of C-reactive protein. Data were also collected regarding any previous or ongoing treatments, including the dose and duration of any corticosteroids, nonbiologic DMARD, biologics, GH, age at biologic therapy onset, number and type of biologics, and the dose and duration of each biologic agent.

**Data analyses.** Height and weight measurements were performed by a pediatrician. Patient height and weight measurements were recorded using the growth chart published by Sempé, et al. Patient target height was calculated using Tanner’s formula. Growth velocity was defined as the change in height SDS during the follow-up. A positive value indicated catchup growth and a negative value indicated impaired growth. Decreased growth velocity was defined as a difference between the height SDS calculated at 2 consecutive measurements that was < −0.5. Height SDS were calculated at 3 different times: at disease onset, at initiation of biologic treatment, and at the latest follow-up. Height SDS corrected for target height was calculated using an auxological data treatment software (KIGS 5). Change in the height SDS before the onset of biologic therapy and during the interval between biologic therapy onset and the latest follow-up were also calculated using reference data for the chronological age from the French population.

Descriptive summary statistics (n, mean, SD, median, interquartile range, minimum, and maximum) were used to summarize continuous variables and percentages for discrete variables. Means were compared using a sample t test, Wilcoxon test, or ANOVA. Categorical data were analyzed using the chi-square test or Fisher’s exact test, in nonpaired data.

Change in the height SDS was compared using paired samples t test. Correlations between change in the height SDS scores and disease-related continuous variables (age at disease onset, disease duration prior to biologic treatments, duration of corticosteroid treatment, or corticosteroid doses in mg/m2) were tested using Spearman’s correlation test. Associations between changes in the height SDS and discrete variables were evaluated using chi-square or Fisher’s exact tests. A univariate and a multivariate analysis of the association between independent variables and change in the height SDS between the onset of the treatment and the end of the follow-up was performed using multivariate linear regression models with a stepwise procedure. For multivariate analysis, we selected all variables associated to change in the height SDS with p value < 0.20 in univariate analysis. All tests were 2-sided. Analyses were performed using R statistical software (www.r-project.org).

**RESULTS**

Among the 780 patients with JIA followed in our center during the 6-month inclusion period, 190 children were taking biologic treatment. Ninety of these patients were not
included: 21 had no followup, 9 had incomplete notes, 7 had another disease that could have affected their growth, 4 were in a wheelchair, 31 had a Tanner stage value of > 3 at biologic treatment initiation, 5 started a biologic treatment for < 6 months, and 13 were taking GH treatment (Figure 1).

At the latest followup, none of the patients had a pubertal delay (defined as no breast development after age of 13 years, or no testis growth after age of 14 years for boys).

The main characteristics of the patients at biologic treatment onset are listed in Table 1. The indications for biologic therapy were active polyarthritis, despite MTX treatment; or severe SJIA or severe uveitis, despite local steroid therapy and MTX, and were in accordance with the practices of the National Reference Center.

The median duration of biologic therapy at the latest followup was 2.92 years (range: 0.59 to 10.17 yrs). Thirty-five patients had < 2 years of biologic therapy at the latest followup and 11 patients had < 1 year. The mean number of biologics per patient was 1.49. Thirty-four patients had received > 1 biologic and 13 of them had received > 2 biologics. Fifty-one patients had received anti-TNF treatment (etanercept, adalimumab, or infliximab); 28 had received an anti-interleukin 1 (IL-1) treatment (7 patients received anakinra first, then canakinumab; 3 patients received only canakinumab); 12 had received the anti–IL-6 receptor antibody tocilizumab; and 9 had received CTLA-4Ig (abatacept). Biologic treatment was combined with MTX in 51 patients. From the onset of biologic treatment to the latest followup, 5 patients had to start steroids. At the latest followup, 45 patients had active disease.

The distribution of height SDS at disease onset, biologic treatment onset, and at the latest followup is shown in Figure 2. All patients had a normal body mass index at inclusion.

The median height SDS of the patients were 0.31 (range –2.47 to 5.46) with a mean of 0.41 ± 1.13. JIA: juvenile idiopathic arthritis; RF: rheumatoid factor; DMARD: disease-modifying antirheumatic drug.

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Uettwiller, et al: JIA, biologics, and growth

Thus, had low growth velocities after the onset of biologic treatment. We analyzed changes in the height SDS (height difference expressed in SDS) between the onset of biologic treatment and the latest followup according to the JIA subtype, steroids usage, the number of biologics, and disease activity at the latest followup (Figure 4). In the multivariate analysis, patients who had been treated with > 1 biologic agent and patients with SJIA had significantly lower changes in the height SDS on biologic treatment than other patients (p = 0.01 and p = 0.04, respectively). Although steroid usage and active disease at the latest followup tended to be associated in univariate analysis (p < 0.20) with lower changes in the height SDS on biologics, none of these associations were statistically significant after adjustment of the other dependent variables.

We also analyzed height SD at the onset of biologic treatment (SD corr TH): 15 patients were below –2SD. After the correction for target height (SD corr TH), 11 patients already had scores below –2SD. Three of them had polyarticular JIA (of whom 1 had uveitis) and 8 had systemic JIA; 9 had received corticosteroids (1 polyarticular and 8 systemics); and 8 had inactive disease and 3 had active disease (all were systemic) at the last followup. These children had been treated with an average of 1.7 biologics and 6 had been treated with > 1 biologic. After the onset of biologic treatment, 4 of these patients maintained low growth velocity up to the latest followup (defined as a ΔSD score of < –0.5; Appendix 1).

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**Figure 2.** Patients’ height distribution. Height distribution is expressed as an SD score (SDS) at disease onset, biologic treatment onset, and the latest followup. Each diagram represents the minimum, 25th percentile, median, 75th percentile, and maximum of the SDS. * Significant difference (p < 0.0001).

**Figure 3.** Growth velocity under biologic treatment. The change in growth velocity from disease onset to biologic treatment onset is shown in relation to the prior growth course; ΔSDS (SD score) represents the growth velocity, expressed as a decelerating growth velocity if ΔSDS < –0.5 or an accelerating growth velocity if ΔSDS > 0.5. While taking biologics, 18 patients still had a decreased growth velocity.

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ΔSDS < 0: 76 patients
Of whom 43 patients with decreased growth velocity
(ΔSDS < -0.5)

ΔSDS ≥ 0: 24 patients

Before biologic treatment onset

On biologics

ΔSDS < 0: 35 patients
Of whom 13 patients with decreased growth velocity
(ΔSDS < -0.5)

ΔSDS ≥ 0: 41 patients
Of whom 12 patients with increased growth velocity
(ΔSDS > 0.5)

ΔSDS < 0: 11 patients
Of whom 5 patients with decreased growth velocity
(ΔSDS < -0.5)
Finally, at the latest followup, 10 patients had a growth retardation (height < –2 SDS) and 1 patient had a height below –2 SDS corrected for target height, but his height was –0.3 SDS not corrected for target height (Table 2). At disease onset, growth retardation < –2 SDS was already documented in 1 patient. Compared to the global population of our study, these 11 patients had the same JIA duration before treatment (median of 1.6 yrs vs 1.8 yrs), the same disease duration (5.4 yrs vs 5.3 yrs), and the same time under biologic treatment (2.8 yrs vs 2.9 yrs), but their disease had started earlier (2.1 yrs vs 4.3 yrs). Eight other children had height SDS that were < –1.5 of their target height at the latest followup. These patients were shorter than genetically programmed, as target height ± 1.5 SD is the generally accepted distribution. The characteristics of these patients are described in Table 2. Nineteen patients (19% of our study group), thus, were below –2 SD or shorter than genetically programmed at the last followup.

**DISCUSSION**

In a cohort of 100 prepubertal or early pubertal children with JIA treated with biologics (but not with GH) and followed in a single reference pediatric rheumatology center, we were able to document a low mean growth velocity at the initiation of biologic therapy and an improvement under biologic treatment. However, in most patients who had developed growth retardation before the onset of biologic therapy, it persisted at the latest followup.

Our cohort of patients had some peculiar characteristics that should be taken into account when interpreting these results. First, a relatively high percentage of the patients had SJIA, a JIA subset that is classically associated with a higher risk of growth failure. In our study, SJIA patients differed significantly from the other patients with respect to their changes in the height SDS under biologic therapy. The larger availability of more active treatment, particularly anti-IL-1 and anti-IL-6 treatments in patients with...
Previous reports have suggested that anti-TNF treatments, the most widely used biologics, have a significant and positive effect on the growth of patients with JIA. A study of a large U.S. cohort of 594 patients treated with the soluble TNF-α receptor etanercept documented a gain of 4.8 percentiles after 3 years of treatment for children below Tanner stage. This height gain, albeit statistically significant, was clinically modest. However, the population was heterogeneous regarding age at biologic treatment onset and some of the children were already pubertal. In another report, 51 of 71 children treated with anti-TNF agents experienced altered growth before starting biologic therapy and these patients gained an average of 0.45 SD after 2 years. Moreover, different subtypes of JIA were followed: 18% of these patients did not respond in terms of height development to etanercept treatment and that was the group of patients with severe and systemic JIA. Two other studies of smaller numbers of patients also indicated similarly encouraging results of anti-TNF treatment in terms of growth velocity improvement and height gain, compared to the pretherapeutic period. Nevertheless, another study of 52 patients treated with etanercept found that prepubertal patients had no significant height gain, despite subgroup analyses that showed a trend toward amelioration. In our study, the high proportions of patients with SJIA and patients with difficult-to-treat JIA may account for the high number of patients who retained a problematic height prognosis. Indeed, our cohort contained 29% SJIA children versus levels of 10% or less in prior studies. In addition, 45% of our children had active disease at the latest followup and a high proportion had been successively treated with several biologics including non-anti-TNF agents. Finally, we included only nonpubertal, prepubertal, or early pubertal children and took into account patient target height, which also increased the proportion of patients in our study who were diagnosed with growth disturbance compared to previous studies. As the great majority of the subjects had no significant pubertal delay, we do not believe that negative value for height SDS could be due to late puberty. Nevertheless, our patients’ growth was better than that reported for patients with JIA who received no biologic treatment, whose final height was below –2 SD in 41% of the patients with SJIA and 11% of the patients with polyarticular JIA.

### Table 2. Patients with growth retardation (height < –2 SD or height correction for target height (TH) < –1.5 SD) at the latest followup.

<table>
<thead>
<tr>
<th>Height at Disease Onset, SD; TH corr SD</th>
<th>Height at Biologic Onset, SD; TH corr SD</th>
<th>Height at the Latest Followup, SD; TH corr SD</th>
<th>Type of JIA (Active = A; Inactive = I)</th>
<th>JIA Duration at Biologic Onset, yrs</th>
<th>JIA Duration at the Latest Followup, yrs</th>
<th>Steroids*, g–²; yrs</th>
<th>Number of Biologics</th>
<th>Age (yrs); Tanner at Last Followup</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.6; –0.5</td>
<td>–2.4; –0.5</td>
<td>–0.5; –0.0</td>
<td>Systemic (A)</td>
<td>6.1</td>
<td>10.2</td>
<td>2</td>
<td>3</td>
<td>11.7; 1</td>
</tr>
<tr>
<td>–1.2; 0.6</td>
<td>–2.6; –1.9</td>
<td>–2.6; –2.1</td>
<td>Systemic (A)</td>
<td>0.8</td>
<td>2</td>
<td>1.7</td>
<td>2</td>
<td>13.8; 5</td>
</tr>
<tr>
<td>–0.6; –1.3</td>
<td>–1.1; –1.8</td>
<td>–3.0; –3.7</td>
<td>Systemic (A)</td>
<td>0.7</td>
<td>5.3</td>
<td>5.2</td>
<td>4</td>
<td>7.4; 1</td>
</tr>
<tr>
<td>1.5; 0.9</td>
<td>–2.2; –2.8</td>
<td>–2.9; –3.5</td>
<td>Systemic (A)</td>
<td>1.9</td>
<td>4.6</td>
<td>19.7; 4.5</td>
<td>1</td>
<td>8.6; 1</td>
</tr>
<tr>
<td>–1.4; –3.3</td>
<td>–1.9; –3.8</td>
<td>–2.8; –4.6</td>
<td>Systemic (A)</td>
<td>4.5</td>
<td>6.9</td>
<td>32.6; 6.8</td>
<td>1</td>
<td>5.4; 1</td>
</tr>
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<td>–0.7; –0.2</td>
<td>–1.3; –0.8</td>
<td>–2.4; –1.9</td>
<td>Polymarticular (I)</td>
<td>1.5</td>
<td>11.3</td>
<td>23.8; 10.7</td>
<td>2</td>
<td>15.1; 1</td>
</tr>
<tr>
<td>–0.4; –1.4</td>
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<td>–2.4; –3.4</td>
<td>Systemic (I)</td>
<td>1.6</td>
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<td>2</td>
<td>6.9; 1</td>
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<td>3</td>
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<td>0.0</td>
<td>1</td>
<td>5.4; 1</td>
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<tr>
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<td>Systemic (A)</td>
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<td>3.4</td>
<td>7.2; 1.9</td>
<td>3</td>
<td>6.3; 1</td>
</tr>
</tbody>
</table>

* The cumulative dose of steroids (g²) was divided by the length of steroids treatment. JIA: juvenile idiopathic arthritis; ERA: enthesitis-related arthritis; RF: rheumatoid factor.
The absence of a significant association between steroid usage and the growth potential of patients taking biologics in our study was in line with a report by Giannini, et al\textsuperscript{16}. In contrast, Tynjäla, et al reported that steroid treatment and disease activity impaired the growth of patients taking biologics\textsuperscript{32}. Steroid treatment may help and hinder the growth of patients with JIA. Its effect in reducing systemic inflammation may be positive in some patients, particularly those with SJIA in whom high systemic inflammation and high IL-6 levels can affect growth\textsuperscript{33,34}. In some children, this effect may counterbalance the well-known negative effect on growth of long-lasting steroid therapy.

Because the growth potential of the patients in our study and in others\textsuperscript{18,35} was already altered before the onset of biologic treatment, earlier treatment with biologics might be the best way to allow patients with JIA to reach their target height. In this regard, however, the median disease duration at biologic treatment onset (16 mos) was relatively short in our study compared to that of previous studies. It remains to be determined whether earlier referral of patients with severe JIA to tertiary care centers or earlier introduction of biologic treatment could improve the growth prognosis of these patients.

Biologic therapy has a positive effect on changes in the height SDS in patients with JIA. However, such therapy might be insufficient to restore normal growth in a subset of patients, particularly in those who do not adequately respond to their first biologic agent. Early initiation of biologic therapy and selecting an optimal first biologic agent may be important options for improving the final height of patients with JIA.

ACKNOWLEDGMENT

We thank Prof. Paul Czernichow for his longstanding help and support in the studies of growth in children with JIA.

REFERENCES


APPENDIX 1. Patients below a –2 SD score (SDS) corrected for target height at onset of biologic. Each line represents the change in the height SDS corrected for target height, per patient, at the onset of disease, at the onset of biologic treatment, and at the last follow-up.