GROWTH HORMONE THERAPY FOLLOW-UP PROGRAMME IN PAEDIATRICS PATIENTS

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Objectives
Adequate evaluation of description and monitoring of pediatric patients treated with growth hormones.

Study design
Retrospective and observational study was performed on patients that initiated growth hormone treatment between years 2009 and 2016. The data we used was obtained from patient's records and collected following the Ministry of Health recommendations:

- **Dose of the drug**
- **Heighth and weight**
- **Weight**
- **Insulin growth (IGF-I)**
- **Value**
- **Growth rate**
- **Bone age**
- **Adult height prediction**
- **Pubertal study**
- **Growth chart**
- **Adverse effects**
- **Adherence to treatment**

Results
32 patients (10±3 years old) 56% children

- **Diagnosis**
  - Classic Growth Hormone Deficiency in 90 %
  - Intrauterine Growth Retardation in 10 %

All of them had data on **height, weight and growth rate** and **growth chart** was not present in any of them.

**Adult height prediction** was only present in 3 % of cases and **pubertal study** in 75 % of cases.

**Bone age** was studied in 87 % of cases and **Insulin growth factor** was determined after 12 months of treatment in 68 % of cases.

**Initial dosage** ✓100%
**Next dosage** ×18%

**Adherence** 90%
**Adverse effects** 12%
- Hyperinsulinaemia, Myalgia and Cephalea.

Conclusions
Treatment monitoring does not comply with stablished criteria. There is a need of **Pharmaceutical Care** in order to guarantee optimal monitoring and security of treatment.

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