Montelukast in early childhood asthma. Predict value of IgG in clinical reply in children 2 to 5 years old?

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ABSTRACT

Background: According to current knowledge, asthma is basically an inflammatory process. Its causes and physiopathological mechanisms are various. The final result is a recurrent obstructive bronchial process, with sibilants and/or dyspnea, which causes an upset in functional respiratory tests, among which the maximum respiratory peak flow meter diminished for the age, sex, and height of patient.

Aims: Our aim is to evaluate if response to treatment with Montelukast has any link with immuno-globulin values (IgG, IgA, IgM, IgE) at start of treatment.

Materials and methods: Included in the study were 32 children, of whom 2 did not begin and 1 who did not provide personal data. There were 29 patients in total, 11 girls and 18 boys. Each made three visits: first where they were instructed, together with their parents, in how to manage the meter and where they received the peak flow meter, Vitalograph, and personal data sheet, where personal and family medical history were noted. The second visit was after 4 weeks, for a clinical assessment and the third visit after 8 weeks. The value register of the PEF would be made morning and night, noting the highest value of three measurements. IgG, IgA, IgM, IgE values were quantified before treatment began. The statistic package STATA 2001 was used in the treatment of data statistics.

Results: Our between the value reached by the PEF after treatment and the IgG values at the beginning of treatment (0.712). In lesser measurement for IgA values (0.660). For each 100 mg/ml of increase in the value of IgG, an increase of 10 l/min in the PEF measurement before and following treatment with Montelukast was produced.

Conclusions: IgG values increase with age. Children with a greater IgG value at the beginning of treatment reached higher PEF values after same. It is not known if the results would be similar with another type of treatment and the way in which IgG influences the results. What appears to be confirmed by available studies is that this relation is found in a group of small children, the aim of our study.

Key words: Asthma. Childhood. IgG. Montelukast.

RESUMEN

Antecedentes: El asma es, fundamentalmente, un proceso inflamatorio, según los actuales conocimientos. Sus causas y sus mecanismos fisiopatológicos son múltiples. El resultado final es un proceso obstructivo bronquial recurrente con sibilancias y/o disnea, que se traduce en una alteración de las pruebas funcionales respiratorias, entre ellas, el pico de flujo
INTRODUCTION

The current concept of asthma involves diverse degrees of clinical expression, as expression of complex histopathological and patho-physiological base, which lead to an inflammatory process in the bronchial tree. In asthma, inflammation is a consequence of the embedded participation of a reticular intricate system in which cells, cytokines, chemical mediators, adhesion molecules, etc. participate in response to different types of aggression, which the bronchial tree suffers.

Due to this knowledge, the treatment of asthmatic disease has adapted toward the control of the different processes involved.

During breast feeding and early infancy, various “asthmatic subgroups” were identified: those which developed sibilants following bronchiolitis, another group which possesses sibilants with a genetical state (“happy wheezing”), usually overweight males and the third group where disturbances in lung function (chronic obstructive pulmonary disease). In a group of very young children, those episodes induced by virus predominated. Therefore, it is reasonable to believe that the immune situation of the child plays a large role in recovery.

In light of current knowledge, within chemical measurements, leukotriens (LT) play an outstanding role. These are fatty acids derived from arachidonic acid which in turn proceed from membrane phospholipids by means of hydrolysis produced by phospholipase A2.

5 Lipoxigenase (5LO), activated by a SLO activating protein (FLAP), acts on the arachidonic acid leading to a series of intermediate unstable compounds which result in the LT primer LTA4, which in turn is also unstable.

In neutrophils, monocytes and alveolar macrophages activated by LTA4 hydrolase, LTB4 is obtained with known chemotactic activity and activator of neutrophils, and to a lesser degree, chemotactic of eosinophils.

In cells fundamentally implicated in inflammation, mastocytes, eosinophils and basophils, LTA4 activated by LTA4 synthase will cause the first of the cysteinil leukotrienes (cys-LT) the LTC4. This undergoes an extra cellular metabolism, which gives rise to LTD4 and LTE4.

The effects of LT are measured by protein G (together with BLT receptor for LTB4 and the receptors of cys-LT): cys-LT1 throughout the entire bronchial tree and cys-LT2 at the venous pulmonary system.

LT produce broncho constriction, exudance of plasma, infiltration of eosinophils, increase in mucous production, contraction of bronchial smooth muscle, disturbance of respiratory flagellum, proliferation of smooth muscle, and specific action as stimulus of liberation of preformed IL-4 in eosinophils; stimulus of the synthesis of eosiinophiotics cytokines (GM-CSF, IL-3, IL-5), increase in expression factor of...
epidermal growth factor, stimulus in production of collagenase by fibroblasts in the lung, etc.

An open prospective study was designed in order to evaluate the efficiency of treatment with Montelukast 4 mg/daily, in the control and functional recovery from bronchial asthma in children between 2 and 5 years of age, attended by consultant paediatricians with primary care, and its possible connection with initial immunoglobulin values (IgG).

METHODS

Children of both sexes with a syndrome diagnoses of asthma were included in this study, who attended their paediatricians at health centers. In this age group, we define as bronchial asthma 3 or more episodes of bronchial obstruction in the last year. Exclusion criteria were as follows: 1) no wish to participate in the study; 2) poor tolerance of medication and/or adverse effects (possible bias); 3) wish to abandon study once started; 4) appearance of another pathology causing suspension of treatment; 5) wish to abandon study once started; 6) appearance of another pathology causing suspension of treatment.

32 patients were originally included. Parents were informed that they could abandon study at any time. One week before treatment began with Montelukast 4 mg (one pill taken nightly but not with meal), the peak flow meter had to be registered morning and night, noting the best of three measurements each time. Instructions were given at the clinic on how to carry out tests using the VITALOGRAPH model (dotted by PROFAS lab) and the form was provided for data. Three programmed consultations were done: inclusion in the study for which parental permission was needed, parents were informed that they could abandon study at any time, as well as how to carry out test and collect variable data. A second consultation was held after 4 weeks and last at 8 weeks. The clinical state of the patient was checked in both, control of peak flow and correct register of data were carried out.

The general variables are: age, sex, height, daily and nightly PEF reading for the 7 days prior to treatment, as well as percentage of immunoglobulins (IgG, IgA, IgM, IgE) (presence and type of adverse reaction, evolutionary clinical markers, need for rescue medication, use of CI, days absent from school).

Statistical analysis

We carried out hypotheses contrasts based on non-parametric statistical tests. To contrast the difference between the proportion of men and women in the sample, an exact binomial queue was used. Hypothesis tests on gender difference for quantitative variables were obtained using a Mann-Whitney test for independent samples. For variables related to evolution, (those measures before and after treatment), a Wilcox test of designated ranges was used for matching samples. In the study for association between quantitative variables and non-parametric coefficient of correlation Spearman was used. As for peak flow, the level increase was measured, while in the rest, decrease was quantified in absolute values. For these indicators, the Spearman coefficient of correlation was constructed with the aim of measuring possible relation of improvement with other variables of interest. Finally, we carried out uni-variance regression models for variables of greater evolutionary interest. Statistic package STATA CORP. 2001.

RESULTS

There exists a greater number of male patients (62.1 %) although the difference is not significant. The contrasts in hypothesis concerning distribution of considered variables comparing both sexes did not reveal any significant difference (significance level of 5%), which suggests that patients have similar characteristics regardless of sex (tables I and II).

Of the 32 patients in the study, 2 did not stand, 1 did not carry out data completion, although the latter did need some variables. 29 patients began the study. 2 patients abandoned study due to adverse reactions (diarrhoea in one, and nightmares and hives in the other). One other patients abandoned study due to a parental decision, 26 patients completed the study overall. These general results are reason for comment in another article.

We find a positive link between all Ig pairs, except between IgG and IgE. We also see an important positive link between IgG and IgA values (coefficient correlation 0.762), therefore the greater level of IgG that exists the greater level of IgA will be found and vice versa.

With regard to the PEF value reached after treatment, there exists a positive correlation with IgG values at the beginning of treatment (0.712), and similarly but to a lesser degree, with IgA values (0.660).

Those children with the highest IgG values at the start of the study, obtained the highest PEF value at the end of the 8-week treatment period with Montelukast. The same occurred in relation to the IgA value, but with a lesser correlation.
No correlation between the IgG value at the beginning of the study and the rest of the considered variables was found. Medication during same (–0.445), as if a greater IgE value meant fewer treatment days, compared with children having lower IgE and requiring. The IgE value was curiously associated although negatively with those days when treatment was required 2 months prior to study (–0.430), and with the need for rescue less rescue medicine during the study (table III).

The adjustment of uni-variant linear regression models revealed a possible link between the IgG value and the increase in PEF, finding no significant statistical link with remaining Ig. This link is maintained even when the multi-variant model is adjusted. For each 100 mg/ml of increase in the IgG value, a rise of 10 l/min is produced in the PEF measurement, prior or following treatment, adjusted for time of evolution, protection rate, IgA and IgM (fig. 1).

As expected, the IgG values increase with age, with the proportion of 7 mg/ml for each month (fig. 2).

### Table I

<table>
<thead>
<tr>
<th>Description of variables according to gender</th>
<th>Male</th>
<th>Female</th>
<th>Total</th>
<th>( p^a )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percentage (%)</td>
<td>62.1 (42.3-79.3)</td>
<td>37.9 (20.7-67.7)</td>
<td>100</td>
<td>0.132</td>
</tr>
<tr>
<td>Age when symptoms began (months)</td>
<td>11.9 (7.6-16.3)</td>
<td>21.0 (10.4-31.8)</td>
<td>15.4 (10.6-20.1)</td>
<td>0.141</td>
</tr>
<tr>
<td>Time of evolution from star of symptoms to treatment</td>
<td>28.9 (20.6-34.8)</td>
<td>23.9 (14.5-33.3)</td>
<td>26.9 (22.0-31.9)</td>
<td>0.233</td>
</tr>
<tr>
<td>Age when treatment began (months)</td>
<td>40.9 (35.5-46.3)</td>
<td>44.9 (37.7-62.5)</td>
<td>42.4 (38.2-46.6)</td>
<td>0.392</td>
</tr>
<tr>
<td>Size</td>
<td>100.9 (87.4-104.5)</td>
<td>103.7 (86.5-110.9)</td>
<td>101.9 (90.7-106.1)</td>
<td>0.486</td>
</tr>
<tr>
<td>Days showing symptoms in last 2 months</td>
<td>5.1 (3.6-7.7)</td>
<td>4.9 (2.1-7.3)</td>
<td>5.0 (3.7-6.4)</td>
<td>0.595</td>
</tr>
<tr>
<td>Treatment days in last 2 months</td>
<td>1.1 (0.2-1.9)</td>
<td>0.8 (0.2-1.4)</td>
<td>1.0 (0.4-1.6)</td>
<td>0.865</td>
</tr>
<tr>
<td>Peak flow prior to treatment</td>
<td>104.7 (73.2-136.2)</td>
<td>92.0 (61.7-122.3)</td>
<td>100.0 (78.5-121.5)</td>
<td>0.649</td>
</tr>
<tr>
<td>Peak flow after treatment</td>
<td>142.6 (109.7-175.6)</td>
<td>125.5 (89.6-161.4)</td>
<td>136.3 (113.0-159.6)</td>
<td>0.256</td>
</tr>
<tr>
<td>Percentage of peak flow prior to treatment</td>
<td>2 (1.0-3.8)</td>
<td>2.5 (1.3-18.5)</td>
<td>2 (2-6)</td>
<td>0.0392</td>
</tr>
<tr>
<td>Percentage of peak flow following to treatment</td>
<td>50 (5.1-55)</td>
<td>26 (5.6-98.8)</td>
<td>43 (7.6-55)</td>
<td>0.104</td>
</tr>
<tr>
<td>Daily variability prior to treatment</td>
<td>32.0 (21.2-42.8)</td>
<td>31.6 (24.5-38.7)</td>
<td>31.8 (24.9-38.8)</td>
<td>0.704</td>
</tr>
<tr>
<td>Daily variability following treatment</td>
<td>16.2 (10.7-21.6)</td>
<td>17.8 (12.7-22.8)</td>
<td>16.8 (13.1-20.4)</td>
<td>0.329</td>
</tr>
<tr>
<td>Variability in the time prior to treatment</td>
<td>34.2 (27.0-41.3)</td>
<td>36.6 (24.1-48.5)</td>
<td>36.0 (29.1-44.8)</td>
<td>0.699</td>
</tr>
<tr>
<td>Variability in the time following treatment</td>
<td>38.6 (28.8-48.0)</td>
<td>36.7 (23.6-49.7)</td>
<td>37.9 (30.6-46.2)</td>
<td>0.812</td>
</tr>
<tr>
<td>Protection index</td>
<td>47.2 (30.0-61.4)</td>
<td>38.2 (27.5-48.8)</td>
<td>43.8 (34.5-53.2)</td>
<td>0.209</td>
</tr>
<tr>
<td>Number of times under medication of asthma attacks during treatment</td>
<td>1.7 (0.3-3.1)</td>
<td>0.4 (0.0-1.0)</td>
<td>1.3 (0.3-2.2)</td>
<td>0.215</td>
</tr>
<tr>
<td>School days lost in quarter before treatment</td>
<td>18.6 (10.3-24.1)</td>
<td>13.9 (7.6-20.1)</td>
<td>17.1 (13.0-21.1)</td>
<td>0.334</td>
</tr>
<tr>
<td>School days lost in quarter following treatment</td>
<td>3.1 (0.9-5.0)</td>
<td>0.1 (0.0-0.3)</td>
<td>2.0 (0.2-3.0)</td>
<td>0.130</td>
</tr>
</tbody>
</table>

*a* Simple average.

*b* p-value of hypothesis contrast.

### Table II

<table>
<thead>
<tr>
<th>Description of variables according to gender</th>
<th>Male</th>
<th>Female</th>
<th>Total</th>
<th>( p^a )</th>
</tr>
</thead>
<tbody>
<tr>
<td>IgG</td>
<td>885.0 (755.4-974.7)</td>
<td>904.7 (753.2-1056.2)</td>
<td>880.3 (797.6-963.0)</td>
<td>0.916</td>
</tr>
<tr>
<td>IgA</td>
<td>107.9 (92.0-133.9)</td>
<td>115.7 (95.3-172.0)</td>
<td>110.9 (98.4-135.3)</td>
<td>0.712</td>
</tr>
<tr>
<td>IgM</td>
<td>121.2 (87.6-154.8)</td>
<td>151.0 (52.0-249.9)</td>
<td>132.6 (93.3-172.0)</td>
<td>0.772</td>
</tr>
<tr>
<td>IgE</td>
<td>143.5 (118.8-268.2)</td>
<td>100.5 (34.9-166.2)</td>
<td>127.6 (48.9-206.3)</td>
<td>0.616</td>
</tr>
</tbody>
</table>

*a* Simple average.

*b* p-value of hypothesis contrast.
DISCUSSION

That Ig values increase with age during development is well known\textsuperscript{12,13}. This involves progressive maturation of the immune system, originating in the foetal period and accelerating from then on, in response to contact with a diversity of environmental agents. There is data which supports the existence of certain differences between boys and girls\textsuperscript{14,15}: significantly higher levels of IgM have been found in women\textsuperscript{14}, and even in some cases IgA in a population aged between 10 and 73\textsuperscript{15}.

Children with higher IgG at the beginning of the study had elevated PEF values at the end, i.e. they showed a better functional respiratory response quantified in an increase of 10 l/min in PFM per each 100 mg/ml which increased the IgG value.

A large variety of results referring to IgG values exists as well as sub-classes of IgG, IgA, etc. and their relationship with asthma or recurrent sibilants in early childhood. It is evident that they play some role in the clinical expression of this pathology in the ages considered, however, this role is still as yet unclear.

From a sample of 37 children, Smith discovered lower levels of IgG sub-classes (IgG1, IgG2, IgG4) in intrinsic and extrinsic asthmatic children, whose proportion was higher in children who had lower initial IgG values\textsuperscript{16}, although a normal concentration of IgG...
in serum does not exclude the possibility of a deficit in sub-classes. Smith suggested the possibility of an etiological role in the respiratory symptoms of some children who showed deficits of the sub-classed IgG mentioned above.

A connection between the IgG sub-class deficiency and bronchitis symptoms in early childhood has recently been found. This is more frequent in these age groups than later, suggesting an increased immaturity of immunity. The etiology of recurrences is often viral in these age groups than in later age groups, meaning that in predisposed children a lower IgG "protective" activity could be involved (neutralization, complementary activation and phagocytosis).

In another recent study of 42 children aged between 9 months and 6 years, lower levels of IgG4 were found in children under 24 months than in the control group. From 2 to 6 years, IgE was found to be increased, while IgG3 and IgG4 (p < 0.05) were lower compared to the controls, suggesting a relationship between lower levels of IgG sub-classes and siblings in early childhood.

In Brazilian children aged 7 to 15, lower IgA levels and sub-classes of IgG were found in the group of severe asthmatics, with an IgG3 deficiency being predominant, although this does not seem to be a reliable predictor for the development of infections in this children.

In Japanese children the prevalence of deficit in IgG sub-classes was 31.6% in breast fed children of 6 to 24 months and 26.7% in controls. There were no significant differences, so the conclusion was reached that sibilants in the breast fed child are not linked to a deficiency in IgG sub-classes, although there exists a link between recurrent siblings and concentration of IgG sub-classes.

In older Asian children with asthma (average 7.5 years), no deficiency in IgG sub-classes was found. Only IgE levels were four times higher than control subjects, suggesting that at these ages there is greater participation of IgE responder as opposed to environmental allergens and a larger maturation of IgG responder.

IgD levels were found to be lower in children with the first signs of atopic asthma (p < 0.001), although these were normal in the course of the following 18 months. Is this a non-specific response or an attempt to block the immune response in asthma favouring tolerance?

The transient absence of IgA in saliva during the first years of life is linked to an increased risk of bronchial hyperactivity BHR, but no link was found between this IgA transient deficiency and the clinical diagnosis of asthma, which would favour the thesis of independence of these two clinical frameworks.

IgA and IgD levels were found to be lower both in patients with intrinsic and extrinsic asthma, compared with the control group. IgA was found quite high in smokers. Women had significantly higher IgA levels compared to men. Both IgG and IgA levels increased with age (14 and 24).

From a sample of 315 apparently healthy males aged between 19 and 22, D’Amelio found no relation between clinical history and the remaining Ig classes, as occurred in our study, where no link between IgE levels and the remaining Ig classes was found in such low age groups as those of our patients.

Those respiratory and non respiratory pathologies where different deficits of IgG sub-classes are found are very diverse and their exact interpretation and comprehension has yet to be fully understood.

In addition, in the broncoalveolar lavage (BAL) in asthmatic patients, the IgG sub-classes were significantly higher than in healthy control subjects, showing a local production of 1 or more sub-classes and an increased exudance of same.

The predominant Ig in BAL is IgA which seems to be produced locally, finding significant statistics for concentration of sIgA and I gA1 patients with allergic rhinitis. The authors suggest the possibility of some role played by this Ig in the mediation of the atopic illness.

When we study the influence of the maternal immune response in a newborn baby and in the early years of life we find some data of interest. The Ac. as opposed to the IgE maternal allergens are consistently linked to the presence of asthma in the mother, while those IgG class Ac. are linked to maternal rhinitis and eczema in the child.

Moreover, treatment with intravenous Ig in patients with severe asthma often leads to clinical improvement, reduction in the need for corticoids and a decrease of cutaneous reactivity in prick-test. Finally, it is important to understand the present situation regarding "hygiene hypothesis" in the development and management of bronchial asthma. Good hygiene in the early years is linked to a lesser prevalence of atopic illness. In a study carried out in Denmark on an adult population and considering seropositivity for Hepatitis A, Helicobacter pylori and Toxoplasma gondii as markers of poor hygiene, a link was found between positivity to 2 of these markers and low prevalence of atopy, while colonization of the intestinal tract by potentially pathogenic bacteria is associated with high prevalence of atopy.

As regards infection by certain pathogens and their relation to atopic illness, new data appears again.

Bjørnsson et al., find a link between recent infection (IgM > 1/16) and/or previous infection (IgG < 1/512 and > 1/32) per Ch1. Pneumonic and the presence of...
The sex of the patient did not influence results. IgG values increase with age being quantified in our case in 7 mg/ml of increase for each month that age increases. The children with the highest IgG values before treatment began reached the highest PEF values after being treated with 4 mg of MONTELUKAST (10 l/min of PEF increase for each 100 mg/ml of increase in IgG value).

CONCLUSIONS

The literature does not clarify the mechanisms implicated in these results. There exist various data which certify the implication of these Ig in the asthmatic process, which seems greater in breast fed and small children. Perhaps a possible explanation could be greater protection as opposed to precipitating factors of asthmatic symptoms, of viral and bacterial type, which their association would have been found significant before. Perhaps a possible explanation could be greater protection as opposed to precipitating factors of asthmatic symptoms, of viral and bacterial type, which their association would have been found significant before.

REFERENCES


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