The Bcl I single nucleotide polymorphism of the human glucocorticoid receptor gene h-GR/NR3C1 promoter in patients with bronchial asthma: pilot study

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Abstract Bcl I in the promoter polymorphism observed within h-GR/NR3C1 gene may play an important role in the development of bronchial asthma and resistance to GCs in the severe bronchial asthma. The aim of the investigation was to study the correlation between this h-GR/NR3C1 gene polymorphism and occurrence of asthma in the population of Polish asthmatics. Peripheral blood was obtained from 70 healthy volunteers and 59 asthma patients. Structuralized anamnesis, spirometry and allergy skin prick tests were performed in all participants. Genotyping was carried out with PCR–RFLP method. In healthy, non-atopic population variants of Bcl I: GG, GC, CC were found with frequency 0.129/0.471/0.400, respectively. In asthma patients Bcl I: GG, GC, CC occurred with respective frequencies of 0.410/0.462/0.128. Chi-square analysis revealed a significantly different (P < 0.05) distribution between cases and controls for the Bcl I polymorphism. The Bcl I polymorphism of h-GR/NR3C1 gene is significantly associated with bronchial asthma, susceptibility to the development of severe form and resistance to GCs in Polish population.

Keywords Glucocorticoid receptor · Glucocorticoid receptor gene polymorphism · SNP (single nucleotide polymorphism) · Inflammation · Resistance to steroids

Introduction

Bronchial asthma is a disease with multifactor etiology [1]. The mutual correlations among the groups of factors predisposing to the development of the disease and the prevalence of asthma are complex in character [2]. The hereditary component of asthma is determined polygenetically [3]. The environmental component is implied significantly by neuroimmune reactions occurring at the molecular level. It should be emphasized that bronchial asthma is a disorder whose primary cause can probably be traced in the disturbed immunoregulatory mechanisms at the lymphocyte level, with secondary overproduction of IgE class antibodies and allergic inflammatory condition [3]. Glucocorticosteroids (GCS) constitute the basic group of medications used to control inflammatory conditions in patients with bronchial asthma. They exert a multidirectional and specific effect on various cell types. They regulate the expression of specific genes within the cell
nuclei via the corticosteroid/receptor complex/receptor
(GCS/GR) [4].

Glucocorticoid resistance is a complex problem [5].
It may be constitutional in character, or develop as a sequel
to an inflammatory process. It should be emphasized that
each tissue represents different sensitivity to GCS. There
are serious doubts whether we are born with steroid-
resistant asthma, or we acquire it during our lifetime, i.e. if
it is dependent on genetic or environmental factors.

The genes involved in increased production of allergen-
specific IgE class antibodies (atopy), bronchial hyperreac-
tivity, production of inflammatory response mediators, and
Th1 and Th2 lymphocyte population sizes play a role in the
etiopathogenesis of the disease [6, 7]. It is notable that the
results of search for the gene or genes predisposing for
the development of atopy or bronchial asthma obtained so far
are not consistent, and the research is still going on.

The human glucocorticoid receptor gene/nuclear recep-
tor subfamily 3, group C, member 1 gene h-GR/NR3C1 is
localized on chromosome 5q31–q32 and consists of nine
exon [8]. The mRNA transcript for h-GR/NR3C1 gene
protein undergoes splicing, which leads to the formation of
four mRNA isoforms: GRα, GRβ, GRδ and GRγ [9, 10].
The only active form of the receptor is GRα. The remaining
isoforms are post-transcriptional modifications of the h-
GR/NR3C1 gene lacking the full potential [3]. The glucocor-
ticoid receptor is a protein made up of a single poly-
peptide chain consisting of 777 amino acids [11]. A few
domains can be distinguished within the receptor. Domain
C, which is the carrier of immunogenicity and other bio-
lological characteristics, is located between amino acids
1–421, which account for a half of the receptor size. The
central GR area (between amino acids 421–486) is
responsible for DNA binding (domain B). It contains cyste-
ine residues forming complexes with zinc, facilitating
DNA binding and determining its tertiary structure. In the
terminal portion of the receptor molecule, there is a seg-
ment (domain A) which controls binding of a hormone
molecule—GCS [12]. The molecular mechanism of action
of GCS involves binding of the specific ligand/glucocor-
ticoid receptor to the sequences of regulator genes encod-
ing the synthesis of anti-inflammatory proteins determining
the clinical effects of GCS.

Resistance to GCS at the molecular level results from
many mechanisms modifying the function of GR in the
cells. Reduced response to drugs of this group can be
explained by decreased expression of GR, impairment of
their ability to bind DNA, or enhanced expression of
transcriptional factors.

Currently, a complex mechanism of GCS resistance
development in patients with severe bronchial asthma is
postulated. Schwartz et al. [13] associated it with various
mechanisms related to GR function, Carmichael et al. [14]
studied the density of complement receptors on monocytes
and demonstrated no changes in GR expression after
administration of GCS, Corrigan et al. [15] described
reduced GR affinity to the ligand as a result of p38MAP
kinase activation by IL-2 and IL-4, Adcock et al. [16]
demonstrated development of GCS resistance associated
with increased AP-1 (activator protein 1) activity, or
increased affinity of AP-1 to GR.

Polymorphisms present within the h-GR/NR3C1 gene
may inhibit formation of GR/GCS complexes, reduce
transcription and cause transrepression of the genes
encoding proteins synthesized within the framework of
cellular response to GCS [17]. Bcl I RFLP is formed as a
result of changes in a single base. A C/G single nucleotide
polymorphism (SNP) within the h-GR/NR3C1 gene pro-
moter has been localized in the intron 647 bp away from
the exon/intron binding site. Allele G is particularly asso-
ciated with sensitivity to GCS. It increases the cellular
response to GCS and occurs less frequently than allele C
[17–19]. Polymorphism Bcl I (C>G) within h-GR/NR3C1
gene promoter demonstrates correlations with sensitivity to
steroids, multiple sclerosis and hypothalamo-pituitary-
adrenal axis [20].

Because of complex etiopathogenesis of bronchial
asthma, its heterogeneous nature, incomplete gene pene-
tration, presence of phenocopies, differentiated gene
expression and gene–gene interactions, thorough investi-
gation of molecular mechanisms leading to the develop-
ment of bronchial asthma is a difficult problem, which
requires continuous evaluation [21].

The aim of the investigation was to study the correlation
between Bcl I single nucleotide polymorphism of h-GR/
NR3C1 gene promoter and occurrence of bronchial asthma
in the Polish population.

Materials and methods

The study was approved by the local ethics committee
(Consent of Research Review Board at the Medical Uni-
versity of Lodz, Poland, No RNN/133/09/KE). At the
commencement of the study, participants were invited to
attend voluntarily. Before admission, written informed
consent was obtained from every patient.

The study was conducted in a group of 55 patients with
bronchial asthma. Asthma diagnosis was established
according to GINA recommendations, based on clinical
asthma symptoms and lung function test. The control arm
included a group of 70 healthy adults, who met the fol-
lowing criteria: no history or symptoms of either bronchial
asthma, or other pulmonary diseases, no history or symp-
toms of allergy, no history or symptoms of atopic derma-
titis, no history or signs of hypersensitivity to aspirin,
negative results of skin tests for 12 common allergens, no first-degree relatives with bronchial asthma or atopic disorders.

Venous blood samples were collected from the participants onto EDTAK3, DNA was obtained from peripheral blood leukocyte fraction. The genetic material was isolated using Wizard DNA Isolation Kit (Promesa, Madison, WI) according to the guidelines provided by the manufacturer. The investigated polymorphisms were analyzed using PCR–RFLP method.

Amplification of DNA segments for BLI polymorphism was conducted using a forward primer (5'-GAG AAA TTC ACC CCT ACC AAC-3') and a reverse primer (5'-AGA GCC CTA TTC TTC AAA CTG-3') according to standard PCR protocol. Starter binding to complementary DNA matrix sites was accomplished at 56°C. Amplified DNA sequences of 418 bp length were obtained. The material was incubated with Bcl I restriction enzyme (New England Biolabs, Neverly, MA) at 50°C for 10 h. DNA fragments containing 263 and 151 bp identified as a set of representative, typical (wild type) alleles were obtained, as well as segments with 418, 263 and 151 bp. RFLP of 418 bp length was identified as the set of polymorphic alleles. RFLP products were separated by electrophoresis on 2% agarose gel, stained with ethidine bromide and observed in UV light. Representative, typical homozygotes, as well as heterozygotes were sequenced and used as internal control.

The obtained results were subjected to descriptive statistical analysis with calculation of arithmetic means and standard deviations. The significance of differences between mean values was determined by means of Chi², P-value. The allel frequencies of BLI polymorphism significantly more frequently. The frequency of allele G versus C correlated with increased occurrence of bronchial asthma in the investigated population (OR = 3.44, CI: 95% confidence interval, CI = 2.03–5.81, χ² = 22.08, P = 2.61e-06). Genotype GG versus CC was characterized by OR = 12.44, CI: 95% confidence interval, CI = 3.87–540.01, χ² = 20.55 with P = 5.82e-06. The carriers of allele G (GG + GC) versus CC developed bronchial asthma significantly more often (OR = 5.44, CI: 95% confidence interval, CI = 2.05–14.41, χ² = 13.16, P = 0.00029). Mutation SNP G/C within the BLI polymorphism of h-GR/NR3C1 gene promoter had a protective effect and reduced the risk of developing bronchial asthma. The occurrence of allele C versus G reduced in a statistically significant manner the risk of the disease (OR = 0.29, CI: 95% confidence interval, CI = 0.17–0.49, χ² = 22.08, P = 2.61e-06). The carriers of allele C (CC + GC) developed bronchial asthma significantly less frequently (OR = 0.08, CI: 95% confidence interval, CI = 0.02–0.25, χ² = 20.55, P = 5.82e-06.)

| Parameter                        | Control group | Group of bronchial asthma patients |
|----------------------------------|---------------|-----------------------------------|
| N                                | 70            | 59                                |
| Gender                           | 34 women:36 men | 42 women:17 men                   |
| Gender (%)                       | 48.6 women:51.4 men | 71.1 women:28.8 men               |
| Mean age                         | 63.11 years   | 50.81 years                       |
| SD for age                       | ±5.00 years   | ±13.65 years                      |
| Min. age                         | 55 years      | 23 years                          |
| Max. age                         | 75 years      | 77 years                          |
| Obliquity                        | 0.19          | -0.32                             |
| Obliquity standard error         | 0.28          | 0.31                              |
| Median                           | 63.00         | 54.00                             |
| Polymorphism of BLI              |               |                                   |
| GG                               | 0.129         | 0.436                             |
| GC                               | 0.471         | 0.455                             |
| CC                               | 0.400         | 0.109                             |

Table 1 Descriptive statistics of the analyzed parameters in controls and cases

The allel frequencies of the investigated polymorphism in both groups demonstrate distribution consistent with Hardy–Weinberg equilibrium [22–24]. Figure 1 illustrates de Finetti distributions of the investigated genotypes [25–27]. Significant statistical differences between the investigated genotypes were demonstrated (P < 0.05). Patients with bronchial asthma presented allele G of the investigated BLI polymorphism significantly more frequently. The frequency of allele G versus C correlated with increased occurrence of bronchial asthma in the investigated population (OR = 5.44, CI: 95% confidence interval, CI = 2.05–14.41, χ² = 13.16, P = 0.00029). Mutation SNP G/C within the BLI polymorphism of h-GR/NR3C1 gene promoter had a protective effect and reduced the risk of developing bronchial asthma. The occurrence of allele C versus G reduced in a statistically significant manner the risk of the disease (OR = 0.29, CI: 95% confidence interval, CI = 0.17–0.49, χ² = 22.08, P = 2.61e-06). The carriers of allele C (CC + GC) developed bronchial asthma significantly less frequently (OR = 0.08, CI: 95% confidence interval, CI = 0.02–0.25, χ² = 20.55, P = 5.82e-06.)

No intra- and inter-group correlations among the gender, occurrence of allergy and age (P > 0.05) were demonstrated.
Discussion

Bronchial asthma is a disease with multifactor etiology. It develops on the background of mutual interactions between genetic and environmental factors. Numerous studies have confirmed that various responses to GCS can be observed in healthy population. The genetic variability with respect to SNPs and mutual gene–gene interactions are the important elements in the development of bronchial asthma. The key loci determining the development of this disease are localized in chromosome 5q31–q32 (gene encoding IL-3, IL-4, IL-5, IL-9, IL-13, GM-CSF, CD14, LTC4S as well as h-GR/NR3C1 and β2-adrenergic receptor).

The tissue specificity of GCS effect on the transcription of genes encoding the synthesis of anti-inflammatory proteins, determining the therapeutic effect, is dependent on many factors. Numerous regulatory elements of DNA within the h-GR/NR3C1 gene promoter, coactivator proteins, cAMP activity, as well as complex protein kinase signal pathways play the crucial role here [3, 28–32]. Binding of the GCS-GR complex with the DNA of regulator gene sequences is a key signal for co-activator proteins CBP (CREB Winding protein) and pCAF (CBP associated factor) as well as SRC-1 (steroid receptor coactivator-1) [1, 3]. Coactivator proteins unwind the DNA strand, combine with RNA polymerase II and bind with the TATA sequence (TATA-box binding protein, TBP). These interaction lead to initiation of transcription [1, 3]. In view of the discussed processes, the interactions between GCS-GR homodimer and regulatory DNA sequences within the h-GR/NR3C1 gene promoter seem to play the key role [3, 17–20]. Clinical observations and studies conducted by many investigators allow to put forward a hypothesis concerning a cause-and-result relationship between the presence of SNPs within the h-GR/NR3C1 gene promoter, and the development of bronchial asthma and resistance to GCS [3, 18–20, 33].

The presented study emphasizes the role of Bcl I polymorphism of h-GR/NR3C1 gene promoter in the etiopathogenesis of bronchial asthma. It should be emphasized that the analysis of SNPs confirmed the correlation between the DNA promoter sequences of the investigated gene and the frequency of bronchial asthma. Undoubtedly, too small sized of the studied groups are still a shortcoming of our project. Nevertheless, the observed variations indicate allele G as a factor promoting the development of bronchial asthma. Mutation G/C within the gene promoter is protective in character and correlates with reduced frequency of the disease. The study needs to be confirmed in large populations of asthmatic patients, especially those with treatment-resistant asthma. It is noteworthy that the significantly increasing prevalence of steroid resistance (35–50%) among patients with severe asthma requires not
Table 2  Descriptive statistics of the analyzed allele of Bcl I polymorphism in controls and cases

| Allele frequencies difference | Heterozygous | Homozygous | Allele positivity | Armitage’s trend test |
|------------------------------|--------------|------------|-------------------|----------------------|
| [G] vs. [C]                  |              |            |                   |                      |
| [GC] vs. [CC]                | [GG] vs. [CC] | OR = 3.44  | CI = [2.03–5.81]  | \( \chi^2 = 22.08 \) | P = 2.613e-06 |
| Allele frequencies difference|              |            |                   |                      |
| [C] vs. [G]                  | [GC] vs. [GG] | OR = 0.29  | CI = [0.17–0.49]  | \( \chi^2 = 22.08 \) | P = 2.613e-06 |

only correlation of SNP frequencies within the regulatory domains of h-GR/NR3C1 gene promoter, but also assessment of gene couplings crucial for the development of asthma and resistance to treatment.

The latest reports pay attention to the role of gene polymorphisms, genetic variations in receptor expressed and extra receptor transcriptional factors in the etiopathogenesis of asthma and steroid resistance phenomenon. Supposedly, disturbances at the level of AP-1 affinity to GR are one of the molecular pathways of cellular resistance to the effect of GCS. Transcriptional factors, and among them AP-1 (activator protein-1) and nuclear factor kappa B (NF-kB) in particular, are the modulators of activity of numerous genes responsible for the development of inflammation, which, binding to GR block GCS-GR binding with DNA sequences of specific response elements (GRE) [34–38].

Conclusions

To date, it has been impossible to define precisely the role of single loci, regions of the particular genes, and individual SNPs of the h-GR/NR3C1 gene in the etiopathogenesis of bronchial asthma and response of the patients to GCS therapy. In view of the applicable paradigms of modern genetics, the key role is attributed to determination of the gene structure and its allelic variants, as well as to assessment of the correlation between the level of expression and biological properties of proteins.

Marker RFLP of the h-GR/NR3C1 gene is an important factor in the development of asthma. In the Authors’ opinion, allele G of Bcl I polymorphism of h-GR/NR3C1 gene promoter demonstrates a marked correlation with the development of bronchial asthma. Its evaluation requires intensive research on hereditary factors linking the particular SNPs with the development of bronchial asthma.

It should be emphasized that no studies concerning the frequency of h-GR/NR3C1 glucocorticoid receptor gene polymorphisms in bronchial asthma patients, including those with severe, treatment-resistant asthma, have been conducted in Poland to date.

The Authors believe that genotyping utilizing the PCR–RFLP techniques will allow in the future to create a bronchial asthma patient profile—phenotype of the Polish patients. The presented study is a preliminary stage for more profound analysis of correlations between the prevalence of the particular allelic variants of the h-GR/NR3C1 gene and the development of bronchial asthma, including severe, treatment-resistant forms of the disease.

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