DNA Methylation Analysis of the Gene CDKN2B in *Gallus gallus* (Chicken)

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Accepted May 15, 2013

The notion of epigenetics includes mechanisms that affect genetic material expression without changing the fundamental sequence of nucleotides in a gene. These mechanisms are essential during the entire development of an organism, primarily in relation to ageing. The epigenome sets the pattern of all epigenetic modifications and is responsible for tissue- and organ-specific cell differentiation by means of selective gene repression and expression (JONES & MARTIENSSEN 2005). The epigenome is dynamic and, unlike the genome, it is different between particular body tissues (EHLRICH 2003). A crucial role in the regulation of gene expression and repression, chromatin modelling and histone modification is played by DNA methylation (CHUANG & JONES 2007; PLACHETKA et al. 2010; GRUBER 2011).

DNA methylation is a biochemical process that controls chromatin structure. The incorporation of a methyl group into cytosine within CpG islands leads to changes in gene transcription activity and chromatin structure and conformation. The area marked with methyl groups is transformed into heterochromatin, and the expression of genes in this area becomes suppressed (REIK et al. 2001; CLEVELAND et al. 2003; OLSZEWSKA 2007).

Molecular biology methods make it possible to analyse the methylation of both the entire genome and particular genes. The basic differentiation is assessed on the basis of quantitative or qualitative 5-methylcytosine assaying (GRYZIŃSKA et al. 2013). The MSP (methylation-specific PCR) technique is a qualitative technique based on the reaction between sodium hydrogen sulphate and nucleic acid (HERMAN et al. 1996; COLLEMAN & RIVENBARK 2006). As a result of the reaction, cytosine or 5-methylcytosine is deaminated in single-strand DNA, and the modified DNA is used as a matrix in the MSP reaction (AZHIKINA & SVERDLOV, 2005). DNA amplification is performed using two pairs of primers containing one or more CpG loci. The primers are designed to differentiate methylated and non-methylated bases. The first pair of primers detects 5-methylcytosine as cytosine. It is thus complementary and joins the unchanged DNA fragment. The other pair of primers is, on the other hand, complementary to the sequence in which 5-methylcytosine has been transformed into thymine (GRYZIŃSKA et al. 2012). The advantages of this method include rapid analysis, the possibility to obtain results from small DNA amounts (5 μg), specificity and sensitivity. Methylation can
be detected even when just 0.1% of the alleles have been methylated (SULEWSKA et al. 2007).

Research on genome methylation or gene-specific methylation has been undertaken chiefly in man to analyse neoplastic transformations. The literature increasingly more often provides results of studies concerning mammalian genome methylation. On the other hand, data relating to the methylation of the avian genome are infrequent. The use of Gallus domesticus for the present study was not arbitrary. Gallus domesticus is a model organism referred to in biomedical and evolutionary research, comparative genomics and epigenetic studies. The publication of the entire chicken genome sequence revealed the existence of synthenic regions between the human and avian genomes which show that despite 300 million years of divergence between mammals and birds, numerous DNA sequences have remained conservative. The genome of Gallus domesticus is the only mapped avian genome and is used as a model in comparative genomics for at least two reasons. First, the genome is relatively small, 1/3 the size of the human genome, yet harbors orthologs to most of the genes in man (GREGORY 2002; ICGSC 2004; MASABANDA et al. 2004; RUBIN et al. 2010).

Polbar chickens are the only autosexing chicken breed in Poland and one of the few autosexing poultry breeds worldwide. The breed stems from the local Greenleg Partridge stock and the paternal Plymouth Rock component. The new breed was named “Polbar”, since an indigenous Polish breed (-pol) was supplemented with the bar gene. The dominant allele (B) inhibits melanin deposition which causes white stripes to appear in the feathers of adult birds. The bar gene is located on the Z chromosome (KAUFMAN 1963; GRYZIŃSKA & NIESPODZIEWAŃSKI 2009). Little Polbar cocks are light grey (occasionally with a touch of regular grey), while little hens are dark, with a black supercilium in the extended line from the eye (Fig. 1a). Only during the first couple of days following hatching is it possible to identify the sex of the chicks on the basis of differential plumage. Afterwards, the sex-specific colour of the down feathers of the little hens and cocks gradually fades away. Mature birds are striped with grey and yellow legs (Fig. 1b). (KAUFMAN 1963; GRYZIŃSKA et al. 2008a, 2008b; GRYZIŃSKA & NIESPODZIEWAŃSKI 2009).

According to available reports, the bar locus contains five genes: micro-RNA 31 (miRNA-31), methylthioadenosine phosphorylase (MTAP), tripartite motif 36 (TRIM36), protein geranylgeranyltransferase type I, β subunit (PGGT1B) and (analysed in this study) gencyclin-dependent kinase inhibitor 2B (CDKN2B). The bar locus genes are associated with melanine activity in melanocytes (DORSHORST & ASHWELL 2009).

The CDKN2B gene encodes proteins similar to one of the two cyclin-dependent kinase inhibitors present in mammals that control CDK4 and CDK6 kinase activity. In human creatinocytes these proteins are induced by TGF-beta cytokine which inhibits cellular growth in the G1 phase of the cellular cycle. Moreover, the CDKN2A/B locus has a key role in cell cycle regulation. It encodes both the ARF protein, which binds the p53-stabilizing protein MDM2, and the INK4 protein, a cyclin-dependent kinase inhibitor. Loss-of-function mutations in CDKN2A are responsible for familiar forms of human melanoma (LIMA et al. 2008; HELSTROM et al. 2010).

The study was aimed at identifying CDKN2B methylation during embryonic development in Polbar chickens by means of the MSP (methylation-specific PCR) technique.

Material and Methods

Animals

The study involved Polbar autosexing chickens. At present, the only Polbar population worldwide is kept at the Felin Laura Kauffman Didactic and Research Station for Small Animals which belongs to the Department of Biological Basis of Animal Production of the University of Life Sciences in Lublin. The population includes 810 hens...
and 80 cocks (GRYZIŃSKA & NIESPODZIEWAŃSKI 2009).

The Polbar breed is covered by a gene pool protection program aimed at the preservation of populations, the maintenance of genetic variability, reduction of the gene share of foreign breeds, as well as the restoration and stabilisation of the phenotypic and genetic parameters of traits typical of local breeds (CALIK 2009; WÓJCIC et al. 2012).

Sample preparation

Female embryos collected on the sixth and eighteenth day of embryonic development comprised the experimental material (10 embryos for each age group). The material was collected in accordance with the 2/B6e/B64 Lublin Local Ethical Commission for Animal Experiments Resolution (Resolution 8/11) dated 15/03/2011.

DNA was isolated from skin cells using a genomic DNA isolation kit (Genomic Midi AX from A&A Biotechnology).

MS PCR

DNA conversion was performed by means of the EZ DNA Methylation-Gold Kit (D5001) from Zymo Research. The obtained eluate was the DNA intended for the PCR reaction. The MSP reaction was conducted in an MJ Research PTC-225 Peltier Thermal Cycler. The reaction primers were designed so that one pair binded at the methylation site, where methylcytosine remained a cytosine, and the other pair at the site where cytosine deaminated into uracil. Table 1 shows the primer sequences used for the PCR reaction.

Investigated gene **CDKN2B** (cyclin-dependent kinase inhibitor 2B) in *Gallus gallus* (chicken). Gene ID: 395076, Location – chromosome Z; Locus NC_006127 (BELLOTT et al. 2010).

### Parameters of the primers for the PCR reaction

<table>
<thead>
<tr>
<th>No</th>
<th>Primer</th>
<th>Sequence 5’-3’</th>
<th>Size (bp)</th>
<th>T_m (C)</th>
<th>T_a (C)</th>
<th>% GC</th>
</tr>
</thead>
<tbody>
<tr>
<td>1A</td>
<td>CDKN2B MF</td>
<td>ATTCGTCGTTTGGAGAGTTGTC</td>
<td>22</td>
<td>61.99</td>
<td>57</td>
<td>41</td>
</tr>
<tr>
<td>1B</td>
<td>CDKN2B MR</td>
<td>CCGTACTAACCGCGCCGCTCTCTACG</td>
<td>21</td>
<td>62.59</td>
<td>57</td>
<td>41</td>
</tr>
<tr>
<td>2A</td>
<td>CDKN2B UF</td>
<td>TGTTGTTGAGAGTTGTTGGG</td>
<td>21</td>
<td>62.53</td>
<td>57</td>
<td>43</td>
</tr>
<tr>
<td>2B</td>
<td>CDKN2B UR</td>
<td>CCATAACTAACCTCTCTACACCA</td>
<td>24</td>
<td>60.84</td>
<td>57</td>
<td>46</td>
</tr>
</tbody>
</table>

T_m – matrix incorporation temperature; T_a – melting temperature; MF – complementary primer for the leading strand at the methylation site; MR – complementary primer for the delayed strand at the methylation site; UF – complementary primer for the leading strand at the non-methylated site; UR – complementary primer for the delayed strand at the non-methylated site.

**Primers**

Primers for MSP were complementary to the gene promoter region. Criteria for the selection of primers: island size >100; GC percent >50.0; obs/exp 0.60 (Fig. 2).

PCR compromised a total reaction volume of 25 μl. A thermal cycler (MJ Research) was programmed for an initial incubation at 94°C for 3 min; 35 cycles each with denaturation at 94°C for 1 min, annealing at 57°C for 1 min, and extension at 72°C for 1 min; and a final cycle at 72°C for 10 min. Amplification products were size-separated by electrophoresis in 1.8% agarose gels and visualized by ultraviolet illumination after staining with ethidium bromide.

**PCR amplification products**

Methylation length (173 bp):
5’ATTTCGTCGTTTGGAGAGTTGTCGGGGTG
GCCGGTTTTGGAGCGCCGATACCGCGCTG
TTTTATGGGTGGITTTTTTCGCGCTGTCG
GGTAGTCGGGTTAGCAGAGGGTTTTC
GGTAGTTATAGATACAGGGTTGTCGGG
TCGGGTATGGGCTAGAGGCGGTTAGTAC
GG 3’

Non-methylation length (173 bp/176bp CCC from the 5’ end)
5’CCCATTTTTTTTTTGGAGAGTTTGGG
GGTGGTTTTTGGAGTTGATATGTGTT
GTTTTATGCTGGTTTTTTTTTTTTTTGGGT
TGGTAGTTGGGGTAGAGGGGTGGTTT
GGTAGTTATAGATAGAGGGGGGGTTGG
TGGGTATTGAGAGTTTGGGTTAGTG
GG 3’

**Table 1**

Parameters of the primers for the PCR reaction
Fig 2. Search result graphic promoters of MSP using MethPrimer.

Fig 3. MSP reaction results for CDKN2B gene on 6th (a) and 18th (b) day of the embryonic development of Polbar chickens (M – methylated cytosine, U – unmethylated cytosine).
Results

The study showed that the CDKN2B was not suppressed on the 6th and 18th day of embryonic development of Polbar chickens. The electropherograms present the results of the MSP reaction (Fig. 3). The intensively luminant MSP product bands for which non-methylated primers were used proved that the amplified product did not undergo methylation.

The complete absence of amplification product using primers overlapping the methylated variant of a given segment of DNA (173bp) on the 6th day of development and the presence of a very weak amplification product on the 18th day of embryonic development may provide a small percentage of methylated cytosines in the analyzed DNA fragment at a later stage of embryonic development.

Discussion

This study aimed at identifying CDKN2B methylation during embryonic development in Polbar chickens by means of the MSP (methylation-specific PCR) technique. We hypothesized that the gene CDKN2B may be methylated during embryonic development (Gryzińska et al. 2013).

DNA methylation is a species- and tissue-specific process. It has been shown that the global methylation level rises as a function of the age of both eukaryotic and prokaryotic organisms (Russo et al. 1996; Bird 2002; Vanyushin 2005; Boks et al. 2009; Lukasik et al. 2009). The DNA methylation level may indicate age, and incorrect DNA methylation can lead to premature ageing (Lukasik et al. 2009). Disorders in DNA methylation control trigger the activation of oncogenes, inactivation of neoplasm suppressor-genes and chromatin stability disorders (Vanyushin et al. 2005; Lukasik et al. 2009).

Methylation is also used to control gene expression in embryogenesis (Plachetka et al. 2010). At the beginning of embryonic development, DNA is almost completely devoid of methyl groups (demethylated). Thus, almost all genes are active. Following organogenesis, a large percentage of genes remain suppressed until death (Ferguson-Smith & Surani 2001; Hake et al. 2004).

Studies of methylation levels during the embryonic development of chickens revealed a statistically significant increase in DNA methylation between the sixth and eighteenth day (Gryzińska et al. 2013).

The experiments of the present study revealed that the CDKN2B gene is not suppressed on the 6th and 18th day of embryonic development of Polbar chickens, despite a significant rise in global methylation (Gryzińska et al. 2013). As one of the five genes responsible for the activity of melanin in melanocytes, the highly active CDKN2B contributes to the production of this pigment that enables sex differentiation in chicks during the first 24 hours following hatching.

The CDKN2B gene in chickens performs a similar function to cyclin-dependent kinase 2A (CDKN2A) inhibitors analysed for other vertebrate species (Kim et al. 2006). Methylthioadenosine phosphorylase (MTAP) and CDKN2A are associated with malignant melanoma (Borg et al. 2000; Behrmann et al. 2003; Goldstein et al. 2008). However, deletions in these genes have also been identified in other types of neoplastic cells (Isii et al. 1999; M’soka et al. 2000). This shows that these genes may play a more significant role in cell proliferation than previously thought.

A plausible mechanism for the sex-related striped pattern in chickens is that the CDKN2A mutation (or mutations) we identified results in premature cell death, which in turn leads to the formation of white stripes lacking melanocytes. This may be then followed by a new wave of melanocytes recruited from a pool of stem cells, which migrate, colonize the feather follicle, produce melanin and form the next black stripe. Thus, the mutations causing sex-related striation in chickens may have an opposite effect than mutations associated with familiar forms of melanoma in humans (Hellström et al. 2010).

Proteins encoded by CDKN2B are the functional equivalents of p16INK4a proteins in mammals (Campos 1991; Dorphorst & Ashwell 2009). INK4A-Arf expression in mice causes melanocytes to age and stimulates the synthesis of the pigment (Sviderskaya et al. 2002).

The CDKN2A gene has also been analysed in reference to the grey colour of the horse coat. The grey coat colour variant in the horse has demonstrated that a cis-acting regulatory mutation in synaptin 17 (STX17) alters expression of nuclear receptor subfamily 4, group A, member 3 (NR4A3) and its downstream target cyclin D2 (CCND2) (Kimbrell Pielberg et al. 2008). The increased expression of CCND2 is thought to increase the rate of melanocyte proliferation and subsequently deplete the population of melanocyte precursors of the hair prematurely while allowing expansive melanocyte growth in the skin. Over the lifespan of the horse, this results in greying of the hair since melanocytes have to undergo cell death as the hair is formed while skin melanocytes durably persist and develop into melanomas. CDKN2B may play a similar role in the striation phenotype of the chicken by altering the melanocyte cell cycle in

IN MANY TYPES OF NEOPLASMS, CHANGES IN DNA METHYLATION ARE THE MOST FREQUENTLY OCCURRING MOLECULAR CHANGES (JABLONSKA & JESIONEK-KLUPICKA 2004). THESE CHANGES MOST OFTEN RESULT IN HYPERMETHYLATION OF PROMOTOR REGIONS LEADING TO ONCOGENE OVEREXPRESSION AND EXCESSIVE CELL PROLIFERATION, HYPERMETHYLATION OF PROMOTOR REGIONS DEBILITATING SUPPRESSOR GENE FUNCTIONS AND HYPERMETHYLATION CAUSING 5-METHYLCYTOSINE TO DEAMINATE INTO THYMINE LEADING TO POINT MUTATIONS IN SUPPRESSOR GENES AND PROTOONCOGENES, AS WELL AS THE INDUCTION OF CHROMOSOME INSTABILITY (LAIRD 1997).

REFERENCES