APPLICATION OF METHYLIGHT APPROACH FOR DETECTION OF CYTOSINE METHYLATION IN FLAX SEEDS GROWN NEAR CHERNOBYL

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INTRODUCTION

When a methyl group binds to a cytosine base, and forms 5-methylcytosine, this process is called DNA methylation, and is an important epigenetic trait involved in various biological processes (He et al., 2011). For proper understanding of epigenetic changes, it is crucial to know the exact location of 5-methylcytosine in the genome. Dynamic changes in DNA methylation during the plant development and stress response are characteristic for promoter region and gene sequences as well (Suzuki and Bird, 2008). Development of the bisulfite modification of genomic DNA represents an important step in expansion of new methodologies in DNA methylation analysis. Bisulfite modification means the DNA alteration where the unmethylated cytosines are converted to uracil while methylated cytosines are detected as unchanged. This modification allows subsequent analysis of PCR products to differentiate methylated and unmethylated cytosines in DNA sequence. Multiple approaches for DNA methylation analysis at the specific locus are based on the bisulfite modification (Clark et al., 2006, Shen and Waterland, 2007).

Analysis of the methylated cytosines in DNA sequence requires the application of relatively cost- and time-consuming techniques. DNA methylation is not transferred during the PCR reaction, therefore the methods for DNA methylation analysis are generally based on modification of genomic DNA using sodium bisulfite (Eads et al., 2000). Initial methods for DNA methylation analysis such as restriction analysis using methylation sensitive enzymes followed by Southern blotting have been replaced with the more advanced approaches in which the genomic DNA is modified using sodium bisulfite. The main reason to develop more reliable and efficient methods for methylation detection was the occurrence of false positive results due to the incomplete restriction digestion and the need of DNA characterized by high concentration and quality (Singer et al., 1990, Eads et al., 2000).

Eads et al. (2000) published a quantitative approach for identification of DNA methylation, so-called MethyLight. The method measures fluorescence real time while combines PCR method and fluorescently labeled TaqMan probes (Eads et al., 2000, Trinh et al., 2001, Olkhov-Mitsel et al., 2014). The MethyLight allows the quantification of methylation at the particular locus in the genome and uses specifically designed oligonucleotides that hybridize to bisulfite modified DNA based on the methylation status in original genomic DNA (Olkhov-Mitsel et al., 2014). The MethyLight is highly sensitive method which allows the detection of methylated alleles even in the presence of a 10000 more unmethylated alleles. This quantitative analysis makes it possible to precisely determine the occurrence of a particular type of DNA methylation (Eads et al., 2000, Hernández et al., 2013). The essential parameters to develop the MethyLight approach are primer and probe design, appropriate selection of fluorescent dye and the optimization of reaction conditions (Olkhov-Mitsel et al., 2014). Currently, this innovative approach is considered as ideal particularly in clinical research because of the very precise identification of methylated cytosines (Zhou et al., 2011).

Redshaw et al. (2014) compared various approaches for DNA methylation analysis: a) MethyLight using quantitative real-time PCR, b) MethyLight using digital PCR, c) restriction digestion using methylation sensitive enzymes followed by quantitative PCR, d) restriction digestion using methylation sensitive enzymes followed by digital PCR and e) PCR amplification of genomic DNA modified using sodium bisulfite with subsequent sequencing of PCR product. The results obtained by MethyLight approach showed more accurate quantification in comparison to the restriction digestion using methylation sensitive enzymes. On the other hand, MethyLight is limited to the analysis of CG dinucleotides which are present in the sequence amplified by specific primers. The aim of the presented study was to apply the MethyLight protocol for quantification of cytosine methylation in the specific gene region. Herein presented study is a part of multi-year investigation of plant adaptation in the radioactive Chernobyl area. In the beginning, two experimental plots have been established near Chernobyl and crops of agricultural importance have been chosen for investigation, e.g. flax, soybean, sunflower, wheat. The majority of experiments have been performed on flax genome and proteome. Thus, flax seeds grown on the experimental fields around Chernobyl were used for analysis. Previous analysis revealed the presence of numerous DNA methylation sites in the genes encoding the fatty acid biosynthesis using the bisulfite sequencing method (Lancikova et al., 2020). Therefore, the MethyLight approach was tested to assess the cytosine methylation in the specific exonic region of Fatty acid desaturase 3A (FAD3A) gene which is involved in biosynthesis of alphalinolenic acid. The applied MethyLight protocol accurately quantified the present methylation without the need of DNA sequencing.

MATERIAL AND METHODS

Experimental plots and flax seed material

Two experimental plots, remediated (non-radioactive) and radioactive and has been established in the close proximity of Chernobyl in 2007. Flax (Linum usitatissimum L.) of Kyivsky genotype has been cultivated on the experimental plots. Flax seeds for the presented study had been harvested in 2013, after six subsequent years of growing near Chernobyl. Flax seeds had been harvested...
from both plots in three biological replicates. Radioactively contaminated plot was located approximately 5 km from Chernobyl Nuclear Power Plant (CNPP), near the village Chistogalovka and soil radioactivity 206Pb ± 1050 Bq kg\(^{-1}\) of 137Cs and 5180 ± 550 Bq kg\(^{-1}\) of 60Sr has been determined. Remediation field was located directly in the Chernobyl town and soil radioactivity 1414 ± 71 Bq kg\(^{-1}\) of 137Cs and 550 ± 55 Bq kg\(^{-1}\) has been estimated. In both experimental plots, contents of aleurite (silt) and pelitic soil ranges from 20 to 30 %. The soils contained 12 % clay, 2 % organic material, and have been characterized as sodic-podzolic with a loamy-sand texture, which is derived from sandy fluvio-glacial deposits. Soil pH of remediated plot was 6.6, and pH of radioactively contaminated plot 5.6. The soil electric conductivity for both plots was 0.20 dS m\(^{-1}\) (Lancíková and Žiarovská, 2020).

DNA extraction

Genomic DNA has been extracted according the method by Rogers and Bendich (1994). Mature flax seeds, 0.5 g (approximately 100 seeds), have been grinded to a fine powder in liquid nitrogen. Extraction solution (50% (v/v) phenol, 0.45 M sucrose, 5mM EDTA, 0.2% (v/v) 2-mercaptoethanol, 50 mM Tris-HCL, pH 8.8) was added to the flax seed powder and shaken for 30 min at 4 °C. Then, proteinase-K has been centrifuged at 10 000 rpm for 10 min for 4 °C, and the DNA solution has been added to an equal volume of chloroform/isoamylalcohol solution (25:24:1) in order to aqueous phase. The aqueous phase was transferred into a new tube. One volume of phenol-chloroform-isooamylocrohol solution (25:24:1) was added to aqueous phase, and mixed vigorously, then centrifuged at 9500 g for 15 min at 4 °C. Extraction using phenol-chloroform has been repeated until clear aqueous phase without brown color. DNA precipitation has been incubated at room temperature for 20 min, and centrifuged at 5000 g for 5 min. Final washing step has been performed using 70% ethanol. DNA pellet has been dissolved in 300 µl TE buffer (pH 8, 10 mM Tris, 1mM EDTA) (Lancíková and Žiarovská, 2020).

Sodium bisulfite modification of genomic DNA

For DNA modification using sodium bisulfite, 100 ng of genomic DNA has been treated with sodium bisulfite using the EpICt Bisulfite Kit (Qiagen) according the manufacturer's instruction. DNA modification protocol has been optimized for low concentration samples, approximately 1 - 500 ng of DNA in a maximum reaction volume of 40 µl has been used. Reaction conditions consisted of a 5 min 95 °C denaturation, 25 min 60 °C incubation, 5 min 95 °C denaturation, 85 min 60 °C incubation, 5 min 95 °C denaturation, 175 min 60 °C incubation, and cool down to 20 °C. DNA degradation has been carried out by sodium bisulfite treatment. Therefore, DNA amount has been significantly reduced after treatment, resulting in average yield 20–30 ng of single stranded DNA (ssDNA) (Lancíková et al., 2020).

Primers and probe design

Quantitative MethyLight approach has been developed for methylation analysis in the particular coding sequence (107 bp, 5’ exon, 3270 – 3671 bp) of Fatty acid desaturase 3A (FAD3A) gene. Primers for PCR amplification of desired gene region were designed as follows: forward primer 5’- TTGGAGGAGGTTGACGATCGTCGATCGAGATTACG-3’ and reverse primer 5’- AATAAAATAACTAAATAACCATCTAATCAATAT-3’. Two fluorescently labeled probes were designed for both variants, methylated and unmethylated gene sequence. The probe sequence for methylated gene region was 5’- CGTGGAGGGTTGACGATCGTCGATCGAGATTACG-3’ and was labeled at the 5’ end using 6-carboxyfluorescenc (FAM, maximum absorbance 494 nm, maximum emission 518 nm) and at the 3’ end using 5-carboxytetramethylrhodamine (TAMRA, maximum absorbance 555 nm, maximum emission 580 nm). The probe sequence for unmethylated variant of gene region was 5’- TGGAGGAGGTTGATCGTATCGTAGATTACG-3’ and was labeled at the 5’ end using Yakima Yellow (analog of VC(TM), maximum absorbance 530 nm, maximum emission 549 nm), at the 3’ end using Black Hole Quencher 1 (BHQI, maximum absorbance 534 nm, maximum emission 549 nm) (Haushalter, 2008). The position of primers and probes with the gene region of interest is shown in the Fig. 1.

Quantitative real-time MethyLight PCR and high resolution melting analysis

Firstly, the positive control (reference sample artificially methylated) has been prepared by CG methyltransferase (M.sso1), (New England BioLabs Inc., Ipswich, MA, USA) and the negative control (reference sample with unmethylated status) was the PCR product of the selected coding region. PCR amplification and subsequent high resolution melting analysis (HRM) has been carried out in a LightCycler® Nano (Roche Applied Sciences, Penzberg, Upper Bavaria, Germany) (Lancíková et al., 2020). Genomic DNA modified by sodium bisulfite was amplified in quantitative real-time MethyLight PCR, the reaction mixture contained 2× Maxima Hot Start PCR Master Mix (2× Hot Start PCR Buffer, Maxima Hot Start Taq DNA polymerase, 0.4 mM of each dNTP, 4 mM MgCl\(_2\), 900 nM of forward and reverse primer, 300 nM of fluorescently labeled probes and 50 ng of modified genomic DNA. The PCR reactions consisted of initial denaturation at 95°C for 10 min, then 45 cycles of denaturation at 95°C for 15 sec and primer annealing at 55°C for 1 min, polymerization step was omitted. High resolution melting analysis has been performed with temperature ramping from 60 to 97 °C at a 0.05 °C s\(^{-1}\) rate (Lancíková et al., 2020). The melting temperature of methylated and unmethylated PCR products was determined. The LightCycler® Nano Software 1.1 was used for data analysis. The reactions were performed in biological triplicates (Lancíková et al., 2020).

RESULTS AND DISCUSSION

Flax seeds grown on the experimental plots near Chernobyl have been investigated for the presence of methylated cytosines using the quantitative real-time MethyLight approach. During the previous investigations, the presence of methylated cytosines has been identified in the flax genes involved in fatty acid biosynthesis pathway, therefore coding sequence of Fatty acid desaturase 3A gene (5’ exon, gene position 3270 – 3617 bp) has been further investigated. Cytosine methylation is a frequent epigenetic modification closely related to the environmental stress. In many cases, DNA methylation can protect the plant genome from further damage caused by stress factor. Epigenetic mechanisms such as DNA methylation can involve alterations in gene expression, changes in chromatin structure and generation of new phenotypes. Moreover, methylation can enhance the plant adaptability to environmental changes and contribute to the generation of stress-resistant phenotype (Thiebaut et al., 2019). Specifically, FAD3A gene methylation can affect the mechanism of de novo fatty acid biosynthesis whereas the gene plays a crucial role in the conversion of linoleic acid to alpha-linolenic acid. In result, this may lower the quality of flaxseed oil. This study was oriented to provide a new methodology for plant DNA methylation research and aimed to test a precise protocol for methylation quantification using the advanced MethyLight method. The certain level of radioactivity is still present in the Chernobyl area which causes serious damage at the genomic level frequently represented by DNA methylation. Therefore, seeds collected from the experimental plots around Chernobyl have been used for analysis. Various techniques are known for identification of DNA methylation. However, here presented real-time quantitative MethyLight approach belongs among the most advanced, straightforward and accurate methods. Aforementioned approach is well established in clinical research and diagnostics even though there are still a little studies dealing with MethyLight in plant sciences. Herein, fluorescently labeled probes were specifically designed for methylated and unmethylated variants of targeted gene sequence. DNA samples extracted from flax seeds harvested from two experimental fields, remediated (non-radioactive) and radioactive, were investigated in biological triplicate. The obtained amplification profiles were compared against known methylated and unmethylated standards to accurately quantify the cytosine methylation. Results showed that targeted gene sequence is fully methylated, at the level 100% methylation, for both variants of flax seeds collected from remediated and radioactively contaminated experimental plots. The amplification profiles for target samples, methylated and unmethylated standards are shown in the Fig. 2. As shown in the fig. 2, target samples correspond to the methylated standard, therefore the complete cytosine methylation in the investigated gene region can be concluded.
Several authors have previously highlighted the most important advantages of MethylLight approach in DNA methylation detection such as the workflow simplicity, multiple samples can be analyzed simultaneously, high sensitivity and efficiency. For instance, this approach requires only low concentration of genomic DNA, there is no need of post-PCR manipulation such as gel electrophoresis which reduces the risk of contamination. On the other side, MethylLight is able to assess cytosine methylation at the single nucleotide, however provides an comprehensive information about the methylation level in specific gene loci (Eads et al., 2000, Ogino et al., 2006, Zhang et al., 2010, Liu et al., 2016). For that reason, the approach such as bisulfite sequencing might be better option for assessment of individual cytosine methylation, however for overall DNA methylation quantification in targeted short gene region, quantitative MethylLight might be a method of choice. The method is the most commonly applied in clinical diagnostics, DNA methylation identification in multiple candidate sequences, assessment of DNA polymorphism, absolute quantification of infrequent DNA methylation using MethylLight droplet digital PCR, gene expression analysis due to the application of fluorescently labeled probes or analysis of repetitive elements (Weisenberger et al., 2005, Zhang et al., 2010, Endo et al., 2015, Yu et al., 2016).

CONCLUSION

In this study, the sensitivity and accuracy of quantitative real-time MethylLight approach for identification of cytosine methylation in flax seeds harvested from Chernobyl area has been tested. Coding sequence of the specific gene, FAD3A, has been investigated for the presence of DNA methylation. The presence of identified cytosine methylations in FAD3A gene might be closely related to the changes in content of alpha-linoleic acid in flax seeds. Therefore, the assessment of accurate method for methylation detection is of great importance. It might be concluded that MethylLight approach is highly suitable for identification of DNA methylation in the particular gene region. It provides reliable and fast results without the need of post-PCR analysis such as gel electrophoresis or sequencing.

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