The Non-coding Mammary Carcinoma Susceptibility Locus, Mcs5c, Regulates Pappa Expression via Age-Specific Chromatin Folding and Allele-Dependent DNA Methylation

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Abstract

In understanding the etiology of breast cancer, the contributions of both genetic and environmental risk factors are further complicated by the impact of breast developmental stage. Specifically, the time period ranging from childhood to young adulthood represents a critical developmental window in a woman’s life when she is more susceptible to environmental hazards that may affect future breast cancer risk. Although the effects of environmental exposures during particular developmental Windows of Susceptibility (WOS) are well documented, the genetic mechanisms governing these interactions are largely unknown. Functional characterization of the Mammary Carcinoma Susceptibility 5c, Mcs5c, congenic rat model of breast cancer at various stages of mammary gland development was conducted to gain insight into the interplay between genetic risk factors and WOS. Using quantitative real-time PCR, chromosome conformation capture, and bisulfite pyrosequencing we have found that Mcs5c acts within the mammary gland to regulate expression of the neighboring gene Pappa during a critical mammary developmental time period in the rat, corresponding to the human young adult WOS. Pappa has been shown to positively regulate the IGF signaling pathway, which is required for proper mammary gland/breast development and is of increasing interest in breast cancer pathogenesis. Mcs5c-mediated regulation of Pappa appears to occur through age-dependent and mammary gland-specific chromatin looping, as well as genotype-dependent CpG island shore methylation. This represents, to our knowledge, the first insight into cellular mechanisms underlying the WOS phenomenon and demonstrates the influence developmental stage can have on risk locus functionality. Additionally, this work represents a novel model for further investigation into how environmental factors, together with genetic factors, modulate breast cancer risk in the context of breast developmental stage.
Author Summary

A woman’s lifetime risk of developing breast cancer is affected by both genetic and environmental risk factors that can be further exacerbated by breast developmental stage. Time periods conferring increased risk are referred to as Windows of Susceptibility (WOS) and, generally speaking, the molecular mechanisms responsible for their effect on breast cancer risk are unknown. Our work presented here on the characterization of the rat Mammary Carcinoma Susceptibility 5c, Mcs5c, locus has identified a region within Mcs5c that interacts with the neighboring gene, Pappa, in an age-dependent manner to influence gene expression via genotype-dependent DNA methylation. Importantly, Mcs5c-mediated gene regulation occurs specifically within a WOS, and these findings represent the first identified molecular mechanisms by which a WOS influences the ability of a locus to affect mammary/breast cancer risk. This work highlights the importance developmental stage can have on genetic risk factor function, and we anticipate that the Mcs5c locus will serve as a model for future studies on WOS in combination with genetic and environmental risk factors.

Introduction

In the United States, breast cancer is the most frequently diagnosed cancer and second leading cause of cancer death among women [1]. Its etiology is complex, consisting of the interaction of both genetic and environmental risk factors whose contribution to overall risk can vary depending on the developmental context of the individual. In general, time periods in which women are more susceptible to initiating events affecting their long term breast cancer risk are broadly referred to as Windows of Susceptibility (WOS) [2]. In humans, the best documentation of a WOS can be found in studies of radiation exposure in women. Women exposed to radiation between 0 and 30 years of age during either the atomic bombings of Japan or for the treatment of Hodgkin’s lymphoma had an increased risk of developing breast cancer later in life compared to women >30 years of age at time of exposure [3,4]. This time period, therefore, represents one of the WOS, and encompasses ages spanning childhood, adolescence, and young adulthood in women. Animals studies performed in rats to model the human WOS phenomenon [5] further suggest the existence of at least two mechanistically distinct susceptibility windows within the larger human WOS, namely, the sexually immature WOS (iWOS) and the adolescent WOS (aWOS). This division of the WOS is most evident in work by Ariazi et al. [6] on a carcinogen-inducible model of breast cancer, where administration to developmentally immature (3 week) and adolescent-aged (7 week) rats resulted in differential carcinoma development depending on age of administration and the carcinogen used. Additionally, although over 80 genetic loci affecting breast cancer susceptibility have been identified in human genome-wide association studies (GWAS) [summarized in 7], their function in relation to developmental stages has not been characterized. In general, while the effects of window specific exposures are well documented, the cellular mechanisms responsible for their function and governing their interactions with environmental and genetic risk factors are poorly understood.

To begin to understand the complex interactions between WOS, genetics, and the environment, we turned to a comparative genomics approach, utilizing a rat model of breast cancer. The rat is an excellent model for this type of study, as not only does its mammary gland and mammary tumor development mimic that of the human condition [8], but, as previously mentioned, it too displays the WOS phenomenon [5,6]. Additionally, inbred rat strains vary in
their susceptibility to carcinogen-induced mammary cancer, allowing for the identification of genetic susceptibility loci through quantitative trait loci (QTL) analysis. This approach was applied in our lab, utilizing the mammary cancer resistant Wistar-Kyoto (WKy) and susceptible Wistar-Furth (WF) inbred rat strains resulting in the identification and subsequent fine-mapping of the Mammary Carcinoma Susceptibility 5c, Mcs5c, locus [9–11]. Mcs5c maps to a 170kb region located in a large gene desert on rat chromosome 5 that shares homology with mice and humans (Fig 1). In both chemical carcinogen and oncogene-induced models of mammary cancer, congenic lines homozygous for the resistant WKy Mcs5c allele showed an approximately 50% reduction in carcinoma number compared to susceptible WF-homozygous controls [11].

Using the Mcs5c locus as a model, we sought to examine the interaction between a genetic risk factor and WOS. We have characterized an 8.5kb temporal control element (TCE) within Mcs5c affecting the expression of neighboring gene Pregnancy-associated plasma protein A, Pappa, in a genotype-dependent manner in mammary epithelial cells (MECs). The function of the Pappa/PAPP-A protein makes it an attractive candidate for involvement in both the WOS phenomenon and breast cancer development. PAPP-A is a protease that acts to positively regulate bioavailability and signaling of the Insulin-like growth factors, IGFs, through the cleavage of IGF binding proteins 2, 4 and 5, IGFBP2/4/5 [12–15]. The specific role of PAPP-A in normal breast development has not been studied, but the IGF-I pathway, in general, is an essential component of breast/mammary gland development, as evident by the severe mammary gland defects of Igf-I and IGF-I receptor (Igf1r) knockout mice [16–18]. The role of IGF-I in breast cancer development is supported by numerous studies which associate the IGF-I signaling pathway with breast cancer initiation and progression [19]. Indeed, in transgenic mice, overexpression of IGF-I in the mammary gland resulted in increased susceptibility and decreased latency to spontaneous and carcinogen-induced mammary adenocarcinomas [20]. Limited studies of PAPP-A function in cancer have demonstrated that increased PAPP-A activity enhanced tumor growth in ovarian and lung cancer cell lines [21,22], and inhibition of its proteolytic function reduced tumor growth in a murine mammary cancer cell line [23].
Furthermore, TCGA data [24] found PAPP-A to be altered in 6% of invasive breast carcinomas, with amplification/mRNA upregulation identified as the most common genetic alterations, and found co-amplification of neighboring loci, encompassing the homologous MCS5C locus, occurring in approximately 1–2% of cases (accessed via www.cbioportal.org; [25,26]).

In this study, we have identified age-specific differences in Mcs5c activity which support the existence of mechanistically distinct susceptibility windows. We have functionally characterized the non-coding Mcs5c locus, finding that it acts during the aWOS to regulate Pappa expression through age-dependent chromatin looping and genotype-dependent DNA methylation. To our knowledge, this study represents the first identification of a molecular mechanism underlying the aWOS phenomenon and highlights the ability of developmental age to influence the activity of a susceptibility locus.

Results

**Mcs5c acts in a mammary gland autonomous manner**

To determine if Mcs5c exerts its effect on carcinoma multiplicity via the mammary gland, transplant experiments were performed. Donor mammary gland tissue from either the Mcs5c resistant 5C-27 line or a Mcs5c susceptible control line was transplanted onto the interscapular fat pad of recipient rats from both genotypes, creating four donor-recipient groups. This direct transplant design allowed for the detection of mammary gland-host interactions and did not result in differential tissue rejection rates, as the lines are isogenic except at the Mcs5c locus. Transplant tissue rejection rates were not statistically significant between transplant groups consisting of donors and recipients with the same genotype versus groups with different genotypes (Chi-squared test, X = .10, df = 1, p-value = 0.75). Results from the mammary gland transplant experiment are shown in Fig 2. Resistant and susceptible rats receiving resistant donor tissue had a transplant site carcinoma incidence of 21% and 27%, respectively (n = 76, 49), while resistant and susceptible rats receiving susceptible tissue had incidences of 42% and 38%, respectively (n = 69, 39). Recipient rats of either genotype that received susceptible donor tissue had higher transplant site carcinoma incidences than those that received tissue from resistant rats. In this way, the carcinoma phenotype was dependent on the donor tissue genotype and was not influenced by the recipient’s genotype, suggesting that Mcs5c acts within the mammary gland. Indeed, logistic regression analysis found a statistically significant donor effect (p-value = 0.0043; recipient effect p-value = 0.825). Thus, it was concluded that Mcs5c acts in a mammary gland autonomous manner to influence carcinoma multiplicity.

**Pappa is differentially expressed in MECs in an age-dependent manner**

Quantitative real-time PCR (qPCR) was used to investigate expression levels of nearby genes in mammary epithelial cells (MECs) of Mcs5c resistant and susceptible rats at 4–12 weeks of age. This age range was chosen as it captures multiple mammary gland developmental windows, including the iWOS (4 weeks), aWOS (6–9 weeks), and adult (12 weeks) time periods. Pappa, located over 517kb away from Mcs5c, was found to be differentially expressed in MECs in an age-dependent manner (Fig 3). In general, Pappa expression levels were dynamic in Mcs5c susceptible MECs during development, while Mcs5c resistant expression remained relatively steady over time. Compared to Mcs5c resistant rats, Pappa expression was increased in susceptible rats by 43% at 6 weeks (Mann-Whitney U test, p-value = 0.015, n = 13 and 15, respectively), 14% at 7 weeks (p-value = 0.05, n = 23 and 19), and 31% at 9 weeks (p-value = 0.0003, n = 23 and 18). Differential expression disappeared by 12 weeks of age (n = 9 and 18), at which point the mammary gland is fully developed and rats are past the aWOS stage [5]. Expression trends were reversed in 4 week old rats, with susceptible animals showing a sharp decrease in
expression relative to resistant rats (p-value = 8e-5, n = 9 and 8, respectively). Mcs5c, therefore, appears to functioning during both the iWOS and aWOS. Unfortunately, we were unable to obtain robust antibodies for analysis of Pappa protein levels in mammary gland tissue. Differential expression in MECs was not observed for neighboring genes Tenascin C, Tnc, and Tumor Necrosis Factor (Ligand) Superfamily, Member 15, Tnfsf15, during the aWOS. However, differential expression of Tnfsf15 was observed in 4 week-old, immature MECs, highlighting the complexity and age-specific nature of Mcs5c locus activity (S1 Fig).

Mcs5c and Pappa physically interact in an age-dependent and tissuespecific manner

Genotype dependent differential expression seen in MECs led to the hypothesis that Mcs5c contained a long-distance acting regulatory element influencing Pappa expression. Such a relationship could be mediated by a physical association between the two regions, resulting in the looping out of intervening DNA sequence. Chromosome conformation capture (3C) was used to identify such an interaction. To create 3C templates, MECs were isolated from the mammary glands of Mcs5c resistant and susceptible animals at 4, 6, 7, and 12 weeks of age. Two fixed bait regions located at the Pappa locus were chosen for extensive analysis of potential interactions with Mcs5c. These regions, P3-1 and P4-1, span approximately 2.4kb and 2kb in size,
respectively, with P3-1 encompassing *Pappa* exon one and a conserved CpG island, and P4-1 falling within the first intron (Fig 4A and 4C). These two regions were chosen for analysis as their degree of sequence conservation suggested that they may be functionally relevant in transcriptional regulation of the *Pappa* gene (Fig 4A). Bait region P3-1 was negative for any interaction with *Mcs5c* at 4, 7, and 12 weeks of age (S2 Fig). Conversely, 3C analysis using bait region P4-1 revealed an 8.5kb region within *Mcs5c* that displayed a high relative interaction frequency (IF) in 6 and 7 week templates, indicative of a physical interaction between the two regions occurring over a distance of 590kb (Fig 4D). 4 and 12 week templates had a much lower IF at this -590 region, leading to the formation of two distinct, age-dependent interaction groups displaying either a strong (6 and 7 week) or weak (4 and 12 week) IF. The difference in IF for these two groups was statistically significant (Mann-Whitney U test, p-value = 1.02e-10, n = 27 and 38 biological replicates, respectively). For all ages, there was no difference in IF between genotypes, indicating that the interaction is age-dependent but not genotype-dependent. We will therefore refer to the -590 looping region of *Mcs5c* as the temporal control element (TCE; chr5:84,428,694–84,437,192; RGSC 5.0/rn5). Three additional *Pappa* bait regions were tested for interactions with the *Mcs5c* TCE at 4 and 6 weeks of age (S2 Fig). Two of these regions,
Fig 4. *Mcs5c* displays an age-dependent and tissue-specific interaction with the *Pappa* gene in MECs. A section of the *Pappa* gene (A) and the full-length *Mcs5c* locus (B) are shown with *BglII* restriction enzyme sites and corresponding primers used for 3C analysis. The TCE is marked by 3C primer E1-2Aa (arrow). Vertebrate conservation from the UCSC Genome Browser (March 2012, rn5) PhastCons track is also shown. (C) The *Mcs5c* locus is shown in relation to the full length *Pappa* gene, with bait regions P3-1 (blue line) and P4-1 (red line) highlighted in the zoomed in image. *BglII* sites and 3C primers are indicated, with an arrow marking the TCE. The green box highlights a conserved CpG island. (D) MEC 3C profile showing the relative interaction frequency (IF, y-axis) between the P4-1 bait region and regions spanning the entire *Mcs5c* locus. The x-axis indicates the distance between the tested region in *Mcs5c* and the P4-1 bait region. The shaded box highlights the temporal control element (TCE) within *Mcs5c* that displayed an age-specific interaction with P4-1, with 6 and 7 week samples showing a strong interaction (dotted lines) and 4 and 12 week samples a negative/weak interaction (solid lines). (E) Colon epithelial cell and liver hepatocyte 3C profiles showed a negative/weak interaction between the TCE (shaded box) and bait region P4-1. The x-axis is the same as in (D), however, not all sites within *Mcs5c* were tested, resulting in a truncated axis. Only *Mcs5c* resistant rats were used for this analysis. For both (D) and (E), each point represents multiple biological and technical replicates, and standard error bars are included. P-values were obtained using the non-parametric Mann-Whitney U test (*, P ≤ 0.05; **, P ≤ 0.01; ***, P ≤ 0.001).

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P4-1A and P4-2, were negative, while the more proximal P3-3 region displayed an aWOS-specific looping interaction that mimicked the TCE/P4-1 interaction. This indicates that the Mcs5c TCE may utilize a more complex looping scheme to facilitate Pappa regulation, and defines the TCE as a functionally important region within Mcs5c. To determine if these interactions are also tissue-specific, 3C profiles were analyzed from 4 and 7 week colon epithelial cells and 7 week liver hepatocytes from Mcs5c resistant rats. The Mcs5c TCE did not interact with P4-1 (Fig 4E) or P3-3 (S2 Fig) in these tissues, implying that the interactions between Pappa and the Mcs5c TCE are tissue-specific in addition to age-dependent. Sequencing of the resistant WKy and susceptible WF TCE alleles revealed 10 variants between the two (S7 Table), and although our 3C results showed that age-specific looping occurs independent of genotype, we speculate that one or more variants may be involved in genotype-dependent expression differences observed during this time period.

The Pappa CpG island shore is differentially methylated in vivo in a genotype-dependent manner

CpG island (CGI) shores are regions located approximately 2kb away from CGIs, and have increasingly been identified as the sites of tissue specific differential methylation associated with gene expression changes [27]. The Pappa looping fragment, P4-1, resides in a CGI shore region (Fig 5A). As this region is a target site of Mcs5c TCE looping, we hypothesized that Mcs5c may affect Pappa expression through an epigenetic mechanism targeted to the P4-1 fragment. Methylation levels for 12 CG dinucleotides within and proximal to P4-1 were examined in MECs of Mcs5c resistant and susceptible rats at 4, 6, 7, 9 and 12 weeks of age using custom designed pyrosequencing assays (Fig 5A). Selection of these timepoints allowed for the

Fig 5. In vivo methylation analysis of the Pappa CGI and CGI shore. (A) The first exon of the Pappa gene is shown in relation to a conserved CGI (green box) and the P4-1 looping fragment (gray box). The location of the 12 shore CGI dinucleotides investigated in this report are indicated and numbered, as are the regions covered by the two pre-made CGI pyrosequencing assays. The CGI assays each examined 5 CG dinucleotides within the island. (B) A scatterplot demonstrating a statistically significant negative correlation between 6 week MEC Pappa expression (x-axis) and shore methylation (y-axis; Pearson correlation coefficient, R = -0.67, n = 18, p-value = 0.0023) is shown. Shore methylation values were obtained by averaging the absolute methylation percentages of the 6 significant shore sites (Sites 1, 3, 6–9) for each individual sample. A linear trend line is shown with the dotted line (slope = -7.88). (C) No correlation was observed between 6 week MEC Pappa expression (x-axis) and CGI methylation (y-axis; Pearson correlation coefficient, R = 0.16, n = 18, p-value = 0.52). CGI methylation values were obtained by averaging the absolute methylation percentages of the 5 sites examined by the CGI-2 assay for each individual sample. A linear trend line is shown with the dotted line (slope = 0.544).

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examination of methylation patterns before, during, and after the aWOS. In general, methylation levels were dynamic across this region, with sites 2–4 consistently displaying the lowest methylation levels (average = 13% methylated) and sites 9–12 displaying the highest levels (average = 68% methylated) (S4 Table). Additionally, there appeared to be few age-specific differences in methylation levels for animals within the aWOS, therefore data for 6, 7, and 9 week old rats were combined within genotypes. Of the 12 sites examined, 6 showed statistically significant genotype-dependent differences in methylation levels after adjusting for multiple comparisons (Mann-Whitney U-test with Bonferroni correction). The percent change in methylation levels along with p-values are shown in Table 1. All statistically significant, genotype-dependent methylation differences occurred during the aWOS and were directionally identical, with methylation levels decreased in Mcs5c susceptible MECs. The percent decrease in methylation levels ranged from 5.0%–22.7%. Additionally, a number of other sites displayed a similar trend, although these differences were not significant after Bonferroni correction. At ages outside of the aWOS, there were no statistically significant genotype-dependent differences in methylation, although sites 1 and 2 displayed a non-significant trend of increased methylation in Mcs5c susceptible MECs at the 4 week time point. We also investigated the methylation state of the Pappa CGI using 2 pre-made pyrosequencing assays (Fig 5A). Methylation levels for both assays were assessed in 4 week old animals, while one assay was examined at the remaining timepoints. For all CGI assays and timepoints, there were no genotype-dependent differences in methylation levels and, in general, the Pappa CGI is hypomethylated at all ages, with site specific methylation levels ranging from 0.16% - 8.41% (S5 Table). The observation of decreased shore methylation and increased Pappa expression in Mcs5c susceptible MECs strongly supports the canonical role of DNA methylation in gene regulation, that is, that the two are negatively correlated. Indeed, for 6 week MECs, for which we had both DNA and RNA samples, Pappa expression was negatively correlated with the average methylation percentage of the 6 significant shore sites (Fig 5B; Pearson correlation coefficient, $R = -0.67$, $n = 18$, $p$-value = 0.0023). By contrast, no correlation was observed between Pappa expression and the average methylation percentage of the CGI-2 assay sites (Fig 5C; Pearson correlation coefficient, $R = 0.16$, $n = 18$, $p$-value = 0.52). The identification of genotype-dependent methylation differences during the aWOS suggests that Mcs5c facilitates genotype-dependent Pappa expression differences observed during this time period through epigenetic modification of the Pappa CGI shore.

Removal of the Mcs5c TCE decreases Pappa expression and increases shore methylation in vitro

In an effort to causally tie the Mcs5c TCE to Pappa expression and CGI shore methylation, the entire 8.5kb region was targeted for deletion in the rat mammary carcinoma cell line, LA7.

### Table 1. Percent change in Mcs5c susceptible MEC methylation at sites within the Pappa CGI shore relative to Mcs5c resistant MECs.

|                    | Site 1  | Site 2  | Site 3  | Site 4  | Site 5  | Site 6  | Site 7  | Site 8  | Site 9  | Site 10 | Site 11 | Site 12 |
|--------------------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| 4 weeks (iWOS)     | +22.6   (NS) | +26.1   (NS) | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    |
| 6/7/9 weeks (aWOS) | -20.4   (0.0002) | -23.1   (NS) | -22.7   (0.0004) | -12.6   (NS) | -5.1    (NS) | -12.2   (0.0004) | -13.4   (0.0003) | -8.9    (0.0114) | -5.0    (0.023) | -5.6    (NS) | -2.2    (NS) | (NS)    |
| 12 weeks (adult MG)| (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    | (NS)    |

The direction of the percent change in Mcs5c susceptible MEC methylation levels relative to Mcs5c resistant MEC levels is indicated by a + (increase) or — (decrease). Bonferroni corrected p-values are shown in parentheses and non-significant percent change trends are also shown (Mann-Whitney U Test; MG—mammary gland; NS—not significant).

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Two CRISPR guides were used to target the region, and clones were screened via PCR across the cut site, with validation by sequencing (S1 Table). We were unable to identify a clone with all copies of the TCE removed, despite much effort. This was likely due to the aneuploid nature of LA7 cells, and mutations incurred at CRISPR guide target regions (S2 & S3 Tables). Copy number analysis of 9 positive CRISPR edited clones showed that we were able to delete a majority of TCE copies, reducing the copy number by 3.5-fold across all clones (Fig 6A). 3C analysis of positive clones indicated that removal of multiple TCE copies resulted in decreased TCE/P4-1 looping, but did not alter TCE/P3-3 looping, which remained consistent with WT levels (Fig 6B). Interestingly, this suggests that the looping mechanisms responsible for these interactions are functionally distinct. Expression analysis revealed a significant reduction in Pappa expression in CRISPR clones compared to wild-type LA7 cells, with Pappa decreased 4-fold across all clones (Fig 6C). A Pearson correlation coefficient was computed to determine the relationship between Pappa expression and TCE copy number, and a positive correlation between the two was observed (Fig 6D; R = 0.6245, n = 13, p-value = 0.0225). Conversely, no change in Tnc or Tnfsf15 expression was observed with TCE knockdown, and expression levels were not correlated to TCE copy number (S3 Fig). These data support our hypothesis that Mcs5c contains a long-range regulatory element, and emphasizes the functionality of the TCE/P4-1 chromatin loop to Pappa gene expression.

Our in vivo analysis highlighted the importance of Pappa CGI shore methylation to Pappa expression, and we sought to verify this relationship in our in vitro model as well. Treatment of wild-type LA7 cells with the DNA methylation inhibitor 5-aza-2'-deoxycytidine (5-aza-dC) resulted in a 15-fold increase in Pappa expression (Fig 6E), indicating that DNA methylation plays a role in Pappa regulation. To more specifically address the relationship between TCE/P4-1 looping and Pappa methylation, CGI and CGI shore methylation levels were analyzed in wild-type LA7 cells and CRISPR edited clones (S10 Table). Two CGI shore sites showed statistically significant differences in methylation, with a 3.9% decrease and a 25% increase in methylation levels observed in CRISPR clones at sites 5 and 12, respectively (S10 Table and Fig 6F). Methylation changes at site 12 were more pronounced, and were investigated further. We found site 12 methylation levels to negatively correlate with TCE copy number (Fig 6G; R = -0.8034, n = 13, p-value = 0.0009). Site 12 methylation levels were also negatively correlated with Pappa expression (Fig 6H; R = -0.6022, n = 17, p-value = 0.011), mimicking the observed in vivo relationship. Altogether, these data suggest a functional chain of events whereby the TCE, via the TCE/P4-1 loop, affects Pappa CGI shore methylation levels which then, in turn, affect Pappa expression.

Discussion

Previous work on Mcs5c had fine-mapped the locus to a 170kb non-coding region on rat chromosome 5. This locus resulted in an approximately 50% decrease in both chemical carcinogen and oncogene-induced mammary carcinoma development when homozygous for the resistant WKy allele. The gene Tnc was initially identified as a possible target of Mcs5c activity, with genotype-dependent differential expression observed in the thymus and ovaries exclusively following carcinogen exposure [11]. However, in this study, we have shown that the Mcs5c locus affects carcinoma multiplicity in a mammary gland autonomous manner (Fig 2). This suggests that the previous non-mammary gland expression differences observed following carcinogen exposure do not play a role in carcinoma initiation, and are either irrelevant or secondary to initial carcinoma development that is dependent on mammary gland intrinsic factors. While these hypotheses warrant further investigation, a reevaluation of gene expression within the mammary gland was conducted, revealing genotype-dependent and age-specific differential
Fig 6. Removal of Mcs5c TCE copies results in decreased Pappa expression in vitro. (A) CRISPR guides targeting the TCE were transfected into LA7 cells, and clones were screened for removal of the target region. Nine positive clones were assessed for remaining TCE copy number via qPCR, standardized to a non-targeted region within the Pappa gene. Copy number in wild-type LA7 cells (n = 4 independent cultures) was also assessed, and results were normalized to diploid MECs. (B) Interaction frequency (IF, y-axis) was calculated in select positive clones (n = 3) and WT LA7 cells between the TCE and Pappa bait regions P4-1 and P3-3. The IF for a positive control region, two nearby BglII fragments, is shown for reference. (C) Pappa expression in positive clones and WT LA7 cells (n = 6) was analyzed via qPCR and standardized to Tbp expression. (D) A scatterplot of Pappa expression and Mcs5c TCE copy number demonstrate a statistically significant positive correlation between the two (Pearson correlation coefficient, R = 0.6245, n = 13, p-value = 0.0225). A linear trend line is shown (slope = 5.327). (E) WT LA7 cells were treated with 0 μM (n = 4) or 1 μM (n = 4) 5-aza-dC for 48hrs. Pappa expression was analyzed via qPCR and standardized to Tbp. (F) Methylation levels at Pappa CGI shore site 12 are shown for WT LA7 cells (n = 8) and CRISPR clones (n = 9). The p-value reflects Bonferroni correction. Scatterplots demonstrating negative correlations for shore site 12 methylation levels and Mcs5c TCE copy number (G) and Pappa expression (H) are shown (R = -0.8034/-0.6022, n = 13/17, p-value = 0.0009/0.011, respectively) along with linear tread lines (slope = -0.046/-0.004, respectively). For all bar graphs, p-values were obtained using the non-parametric Mann-Whitney U test, and standard error bars are shown (*, P ≤ 0.05; **, P ≤ 0.01; ***, P ≤ 0.001).
expression of \textit{Pappa} in mammary epithelial cells (MECs; Fig 3). Specifically, \textit{Mcs5c} susceptible MECs from 6 to 9 week old rats showed increased expression of \textit{Pappa} compared to \textit{Mcs5c} resistant rats. Importantly, the 6 to 9 week age range encompasses a time period of rapid mammary gland development and maturity, falling within the aWOS [5].

We hypothesized that \textit{Pappa} expression changes were mediated by a regulatory element within \textit{Mcs5c}, and our experimental results support this hypothesis, identifying a complex set of mechanisms underlying \textit{Mcs5c}-mediated regulation of \textit{Pappa}. Through 3C experiments, we have identified a region within \textit{Mcs5c}, the temporal control element (TCE), that physically interacts with the \textit{Pappa} locus at two regions, P4-1 and P3-3, in an aWOS- and MEC-specific manner over distances of 590kb and 580kb, respectively (Fig 4 and S2 Fig). The importance of the TCE/P4-1 long-range looping interaction to \textit{Pappa} expression was demonstrated in vitro, where removal of TCE copies resulted in a reduction of TCE/P4-1, but not TCE/P3-3, looping, and correlated with decreased \textit{Pappa} expression (Fig 6). These data indicate that the two observed TCE chromatin interactions are functionally distinct, and demonstrates a strong positive regulatory relationship between the \textit{Mcs5c} TCE and \textit{Pappa} expression, which appears to be dependent on TCE/P4-1 chromatin looping. The \textit{Mcs5c} TCE/Pappa P4-1 interaction, therefore, represents another example of a long-distance acting regulatory region, akin to those identified for the \textit{Shh} [28] and \textit{Sox9} [29] genes, as well as the previously characterized \textit{Mcs1a} locus [30].

Importantly, as looping occurs in a genotype-independent manner, additional mechanisms must be responsible for the differential expression observed between \textit{Mcs5c} resistant and susceptible rats. With the intronic P4-1 looping region falling in a CGI shore, DNA methylation of this region became a mechanistic candidate to explain observed expression differences. The importance of differentially methylated CGI shores to gene expression was first highlighted by Irizarry and colleagues in 2009 [27]. Since then, many studies have shown an association between differentially methylated shore regions and gene expression changes [31–38]. Our study identified 6 CG dinucleotides within and proximal to the P4-1 looping region that were differentially methylated between \textit{Mcs5c} resistant and susceptible MECs (Table 1). Significant methylation differences were observed exclusively during the aWOS, and a negative correlation between shore methylation levels and \textit{Pappa} expression strongly suggest that DNA methylation plays a role in differential \textit{Pappa} expression (Fig 5B). This correlation was recapitulated in our in vitro model, where differential shore methylation was also negatively correlated with TCE copy number (Fig 6G and 6H). As copy number acts as an indicator of TCE/P4-1 looping frequency in this model, this suggests a functional relationship between looping and shore methylation, where the TCE/P4-1 loop acts to facilitate differential methylation that, in turn, regulates \textit{Pappa} expression. Given the inherent difficulties of modeling an age-dependent phenomenon in vitro, these results must be interpreted cautiously; however, we feel that the similarities between our in vitro and in vivo results indicate that these mechanisms are robust and functionally relevant to \textit{Mcs5c}-mediated \textit{Pappa} regulation.

Overall, we have identified two mechanisms associated with \textit{Mcs5c} regulation of \textit{Pappa} expression during the aWOS, chromatin looping and DNA methylation. Our in vitro experiments have indicated the importance of the TCE/P4-1 loop for \textit{Pappa} expression and shore methylation; however, in vivo analyses have shown that these actions are mechanistically distinct, as \textit{Pappa} expression and differential methylation, but not looping, are genotype-dependent. An unresolved issue is precisely how \textit{Mcs5c} is mediating these activities. We hypothesize that the genotype-independent TCE/P4-1 loop serves to facilitate the recruitment of transcription factors, cofactors, and/or methyltransferases that act separately or together to directly regulate \textit{Pappa} methylation and expression during the aWOS. The binding of these regulatory factors would be affected by one or more variants within the TCE without affecting chromatin
looping. Wright et al. [39] identified a similar interaction at the c-MYC locus, where an enhancer-associated SNP affected transcription factor binding without altering chromatin structure. Sequencing of the resistant WKy and susceptible WF Mcs5c TCE alleles (chr5:84,428,694–84,437,192; RGSC 5.0/rn5) has revealed 10 candidate polymorphisms (S7 Table) for future investigation of their effect on protein binding and subsequent Pappa expression and methylation changes.

Mcs5c activity during the aWOS stands in stark contrast to that observed during the iWOS (4 weeks). Specifically, differential Pappa expression during the iWOS is reversed compared to the aWOS (Fig 2), TCE/Pappa looping is lacking (Fig 4D and S2 Fig), and there are no statistically significant CGI shore methylation differences (Table 1). These data indicate that the regulatory actions of Mcs5c are dependent on developmental context, a phenomenon observed at other regulatory regions, most notably the β-globin locus control region [40]. Age-specific differences in Pappa expression, looping, and methylation could be explained by interactions with proteins specific to these developmental time points. Identifying proteomic differences between the immature and adolescent mammary gland will be crucial in understanding the players driving window-specific mechanistic differences in Mcs5c activity. We hypothesize that age-specific protein expression results in an alternative looping interaction between Mcs5c and Pappa during the iWOS. Differential expression of Tnfsf15 exclusively at the 4 week time point (S1 Fig) indicates that Mcs5c may exhibit a more complex chromatin interaction during the iWOS, regulating multiple genes simultaneously. Additionally, a trend towards increased methylation in Mcs5c susceptible MECs is functionally consistent with the reduction of Pappa expression observed during this time period. It is possible that these sites are indicative of significant methylation differences occurring at sites not examined in this study, both at the Pappa locus as well as Tnfsf15, and shore methylation may still, therefore, be relevant to Mcs5c activity during the iWOS.

Altogether, we have functionally characterized the Mcs5c locus, finding that it acts via two distinct mechanisms to influence Pappa expression in an age-dependent manner during a well-characterized breast cancer WOS (Fig 7). This work highlights the importance of characterizing genetic risk factors in the context of developmental windows of susceptibility (G x WOS), and emphasizes the complex interaction between genetic, environmental, and age-specific risk factors. Mcs5c susceptible rats showed increased expression of Pappa in MECs and an increased susceptibility to carcinogen-induced mammary carcinogenesis, supporting a protective benefit of reduced Pappa levels during adolescent development. Decreased levels of Pappa in the developing mammary gland would result in reduced Igf-I bioavailability through a reduction in Igfbp cleavage [12]. Given that the Igf-I signaling pathway acts to promote proliferation and inhibit apoptosis during mammary gland development [41], it is therefore likely that a reduction in free Igf-I would reduce the proliferative index of MECs. As the effects of many environmental mutagens, such as radiation and chemical carcinogens, are dependent on interactions with the DNA of proliferating cells [42], this would result in fewer targets for mutagenesis, and represents one possible method by which reduced Pappa expression during the aWOS may result in a mammary carcinoma resistant phenotype. Understanding the mechanisms behind G x WOS interactions and how environmental risk factors influence these interactions will play a crucial role in breast cancer risk assessment, and in the identification of targets and strategies for cancer prevention in young women. There is growing concern over the impact adolescent exposure to a broad range of environmental factors may have on long-term breast cancer risk [43]. Our work has demonstrated a functional relationship between genetic risk factor activity and developmental stage, and it is likely that environmental risk factors may further confound such interactions. Indeed, CGI shore methylation has been found to be affected by environmental factors such as ELS and diet [32,33,44,45]. We believe that the
Mcs5c locus will serve as a robust model to study how environmental factors affect breast cancer risk by influencing G x WOS interactions, and may encourage the characterization of other such cancer susceptibility loci in this context.

Materials and Methods

Animals

Congenic rat lines were maintained in an AAALAC-accredited facility as previously described [11]. All protocols were approved by the University of Wisconsin–Madison School of Medicine and Public Health Animal Care and Use Committee. Congenic rat lines are defined as having the resistant Wistar-Kyoto (WKy) Mcs5c allele introgressed on a susceptible Wistar-Furth (WF) background. The resistant congenic line used in this study, 5C-27, is WKy-homozygous for a genomic region that includes the entirety of the Mcs5c locus (Fig 1) [11]. Susceptible control animals are WF-homozygous at the Mcs5c locus.

Mammary gland transplantation

Mcs5c WKy-homozygous congenic rats from line 5C-27 were used as resistant donors and recipients (Mcs5c resistant), and Mcs5c WF-homozygous rats were used as susceptible controls (Mcs5c susceptible). Abdominal and inguinal mammary glands were collected from female
donor rats aged 30–35 days old, scissor minced, and split into four equal volumes. One volume was then grafted onto the interscapular white fat pad of four different 30–35 day old female recipient rats. Three weeks after transplantation (51–56 days), recipients were administered the chemical carcinogen 7,12-dimethylbenz(a)anthracene (DMBA), as a single oral dose dissolved in sesame oil at 65 mg/kg of body weight to induce mammary carcinoma formation. At 15 weeks post-DMBA, animals were removed from the study and carcinomas present at the transplant site greater than 3x3mm were counted. Generally, recipient rats developed ≤1 carcinoma at the transplant site, so incidence values were used as opposed to multiplicity. Fat pads were whole mounted and stained with aluminum carmine to verify transplant mammary gland growth. Four transplant groups were studied, with resistant 5C-27 or susceptible donor glands transplanted into both resistant and susceptible recipients (R- > R, R- > S, S- > R, S- > S). The tissue rejection rate for transplant groups consisting of donors and recipients with the same genotype (R- > R, S- > S) was compared to the rejection rate for groups with differing donor and recipient genotypes (R- > S, S- > R) via a Chi-squared test. The effect of donor and recipient genotype on carcinoma incidence, converted to a binary response value, was analyzed using logistic regression with two independent variables (donor genotype and recipient genotype) and no interaction term.

Tissue collection and DNA/RNA processing

For all experiments, mammary epithelial cell (MEC) isolation began with fresh mammary glands (abdominal and inguinal, with lymph nodes removed) that were finely minced and digested for 2 hours at 37°C in 10 mL of Gibco Dulbecco’s modified Eagle’s medium/F12 (DMEM/F12; Life Technologies) containing 0.01 g/mL of type III collagenase (Worthington). Cell pellets were collected by centrifugation and resuspended in 5 mL DMEM/F12. The suspension was loaded onto a 40μm nylon filter to eliminate stromal cells and collect mammary ductal fragments, consisting of an enriched MEC population. DNA was isolated from cells via the DNeasy Blood and Tissue kit (Qiagen). To isolate RNA, cells were homogenized in TRI Reagent (Ambion), followed by RNA extraction using the MagMAX-96 for Microarrays Total RNA kit (Ambion). LA7 cells used for downstream analysis were collected via treatment with 0.25% trypsin/EDTA (Life Technologies). RNA was extracted using the RNeasy Mini Kit (Qiagen) and DNA was extracted using the DNeasy Blood and Tissue kit (Qiagen).

Quantitative real-time PCR

MECs were collected at 4–12 weeks of age from female Mcs5c resistant and susceptible control rats. RNA was isolated as described above. For in vivo and in vitro expression analysis, cDNA was prepared from 1–2μg total RNA using Superscript II reverse transcriptase (Invitrogen). Gene expression was quantified using pre-designed or custom made TaqMan qPCR assays (Pappa, Rn01458295_m1, FAM; Tbp, Rn01455646_m1, VIC; Tnc, probe-FAM 5’ CGAGAGC TGTGATTAGA 3’, primers 5’ GGCTGTCAGAAGGCCAGATG 3’ and 5’ TGCCATGAA GGGATTTGAAGA 3’; Tnfsf15, Rn00595596_m1, FAM) and run on an ABI Prism 7900HT (Applied Biosystems). Tbp was chosen as the reference gene as its expression has been found to be relatively stable across a variety of rat tissues and during different stages of the estrous cycle in the mammary gland [46]. cDNA was diluted 1:4 or 1:8 and run using reaction conditions described previously [11]. Transcript quantities were calculated as described in Smits et al. [30], using a standard curve method to calculate Ct values and extrapolate quantity values. Sample measurements are an average of 3–4 technical replicates and data were analyzed using SDS software version 2.2.2 (Applied Biosystems).
Chromosome conformation capture (3C) assay

Sample templates were prepared from MECs, colon epithelial cells, liver hepatocytes, and LA7 cells. MECs were isolated from 4, 6, 7, and 12 week old Mcs5c resistant and susceptible rats and the resulting cell suspension was diluted in PBS and fixed via the addition of 1.5% formaldehyde. Colon epithelial cells were isolated from 4 and 7 week old resistant rats, processed as described in Whitehead et al. [47], and fixed in formaldehyde. To isolate hepatocytes, the livers of 7 week old resistant rats were digested via cannulation of the portal vein and blanching of the liver using a pre-warmed solution of HBSS (Gibco) + 0.5mM EGTA followed by digestion via pre-warmed DMEM-low glucose (Gibco) + 1000CDU/mL Collagenase type IV (Worthington). Digested livers were collected in DMEM/F12 + 10% FBS on ice and cells dispersed manually. The suspension was filtered through a 100μm nylon filter and the filtrate spun for 2 minutes at 50xg. Supernatant was removed and cell pellets were washed until media became clear, followed by fixation in formaldehyde. Bacterial artificial chromosomes (BACs) encompassing the rat Mcs5c and Pappa promoter regions (CH230-433D12, CH230-498D4, CH230-256M9, and CH230-244C7) were ordered from Children’s Hospital Oakland Research Institute (CHORI) and used as positive control templates. Subsequent template preparation for all cell types and for BAC controls continued as described in detail in Smits et al. [48]. The restriction enzyme used was BglII. Chromatin interactions were detected via PCR, with bait primers located at the Pappa gene tested against Mcs5c primers spanning the entire locus (Fig 4A and 4B). Primer sequences are listed in S6 Table. Reaction components were 1X Herculase reaction buffer, 0.2mM dNTPs, 0.4μM primers, 0.3μL Herculase Enhanced polymerase (5U/μL, Agilent) in a total volume of 25μL. The amount of DNA template to add and optimal annealing temperatures were determined empirically. PCR reactions were performed using the following cycling conditions: 95°C for 1 min, 36 cycles of 95°C for 30 s, Tb for 30 s, 72°C for 20 s, followed by a final extension of 72°C for 8 min. Reactions were analyzed by agarose gel electrophoresis and visualized by ethidium bromide staining. Band intensities were quantified using ImageQuant software (GE Healthcare). A relative interaction frequency (IF) was calculated by dividing the band intensity of the sample templates by that of the BAC control.

Sequencing

Sequencing of the 8.5kb Mcs5c looping region (TCE; chr5:84,428,694–84,437,192; RGSC 5.0/ rn5) identified in 3C experiments was performed on MEC DNA from Mcs5c resistant and susceptible rats to assess polymorphisms between the WKy and WF alleles. Sequencing primers are listed in S7 Table. Traditional Sanger sequencing was performed at the University of Wisconsin–Madison Biotechnology Center DNA sequencing facility as described in Smits et al. [30].

Cell culture and in vitro CRISPR editing

The rat mammary carcinoma cell line, LA7, was obtained from the American Type Culture Collection and maintained in DMEM/F12 supplemented with 100 IU/mL penicillin, 100 μg/mL streptomycin (Life Technologies), 5% FBS (HyClone), and 0.005mg/mL insulin. Gene expression analysis proceeded as described above, and copy number analysis was performed via SYBR Green qPCR (Life Technologies). For 5-aza-2’-deoxycytidine (5-aza-dC; Sigma) experiments, cells were treated with 1μM 5-aza-dC for 48hrs followed by cell collection and processing. For quantification of Mcs5c TCE copies, a primer set located within the CRISPR targeted region was used (5’ CAATCACGTTCACTGTGGGT 3’ and 5’ TCACCTCACACTCCCCATG 3’) and as a control region, a primer set located within the non-targeted Pappa gene was used (5’ TCCTCCTGACCACCTTGGAGA 3’ and 5’ CCCTACAAACAGCAGAGGGAGA
3”). The CRISPR-Cas9 plasmid pSpCas9(BB)-2A-Puro (PX459) was provided by Dr. Feng Zhang (Addgene plasmid #48139) [49]. Guide sequences were designed using the CRISPR Design Tool (http://crispr.mit.edu) and phosphorylated and annealed guide oligos were inserted into the PX459 plasmid via a combination digestion/ligation reaction. 100ng PX459 plasmid was mixed with 2μL of oligos (diluted 1:250), 1μL Fast Digest BbsI (Thermo Scientific), 1X Fast Digest Buffer, 1mM DTT, 1mM ATP, and 1500 units T7 ligase (New England BioLabs) and incubated in a thermocycler for 5 minutes at 37°C followed by 5 minutes at 23°C for a total of 6 cycles. The resulting reaction was then treated with Exonuclease V (NEB) according to the manufacturer’s protocol. The product was transformed into competent cells, and colonies expanded and verified via sequencing of the guide insertion site. LA7 cells were transfected with two CRISPR guides flanking the 8.5kb target region. Transfection was performed via electroporation using a Nucleofector II Device and Amaxa Cell Line Nucleofector Kit V (Lonza), according to the manufacturer’s instructions. Stable clones were isolated following puromycin selection, and clonal colonies were expanded. Removal of the targeted region was determined via PCR screening and sequencing. Primers used to create guides and screen clones are listed in S8 Table.

DNA methylation analysis of the Pappa gene

DNA was isolated from wild-type LA7 and CRISPR edited cells as well as Mcs5c resistant and susceptible MECs at 4, 6, 7, 9, and 12 weeks of age. Bisulfite conversion was carried out on 500ng of DNA using the EZ DNA Methylation-Lightening kit (Zymo Research), according to the manufacturer’s instructions. Four primer sets were designed to amplify the 12 CpG sites of interest within the Pappa CpG island (CGI) shore. Their sequences, along with the sequencing primers used for pyrosequencing, are listed in S9 Table. Optimal amounts of template DNA, MgCl2, primers, and annealing temperature were experimentally determined (S9 Table). In general, PCR reactions were performed using the following cycling conditions: 95°C for 5 min, 50 cycles of 95°C for 15 s, Ta for 30 s, 72°C for 30 s, followed by a final extension of 72°C for 5 min. 15μL of PCR product was used for pyrosequencing on a PyroMark MD instrument (Qiagen), with 2–3 technical replicates per sample. Data were analyzed using PyroMark CpG software (v 1.0; Qiagen). For analysis of the Pappa CGI, 2 pre-made assays were obtained from Qiagen (CGI-1, Rn_D3ZNQ7_01_PM; CGI-2, Rn_D3ZNQ7_02_PM), with PCR conditions following manufacturer’s recommendations. Both pre-made assay CGI-1 and CGI-2 amplified 5 CpG sites within the Pappa CGI, for a total of 10 sites in the island examined. For statistical analysis of methylation differences, the non-parametric Mann-Whitney U test was used, with a Bonferroni correction applied for multiple comparisons.

Supporting Information

S1 Fig. Tnc and Tnfsf15 expression in mammary epithelial cells. Tnc (A) and Tnfsf15 (B) expression was examined in MECs of Mcs5c susceptible and Mcs5c resistant rats at various ages. Gene expression relative to Mcs5c susceptible levels was determined via qPCR with Tnc and Tnfsf15 standardized to Tbp expression. There were an average of n = 12 animals per group, and p-values were obtained using the non-parametric Mann-Whitney U test. Standard error bars are shown (*, P ≤ 0.05; **, P ≤ 0.01; ***, P ≤ 0.001).

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S2 Fig. Chromosome conformation capture profiles for additional Pappa bait regions. (A) A full 3C profile shows the relative interaction frequency (IF, y-axis) between the bait region P3-1 and regions spanning the entire Mcs5c locus in MECs at various ages. The x-axis indicates...
the distance between the tested region and P3-1 (UCSC Genome Browser, March 2012, rn5). (B) Three additional Pappa bait regions (P3-3, P4-1A, and P4-2; see Fig 4A for genomic locations) were tested for interactions with the Mcs5c TCE in MECs from 4 and 6 week old rats. In both (A) and (B), Mcs5c genotypes were combined within time points. (C) The Pappa bait region P3-3 was tested for interaction with the TCE in 4 and 7 week colon epithelial cells and 7 week liver hepatocytes. Only Mcs5c resistant rats were used in this analysis. For all graphs, multiple biological and technical replicates were used, and standard error bars are shown. P-values were obtained using the non-parametric Mann-Whitney U test (MEC—mammary epithelial cells; *, P ≤ 0.05; **, P ≤ 0.01; ***, P ≤ 0.001).

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**S3 Fig. Tnc and Tnfsf15 expression is unaffected by TCE knockdown.** (A) Tnc and Tnfsf15 expression in positive clones (n = 9) and WT LA7 cells (n = 3 independent cultures) was analyzed via qPCR and standardized to Tbp expression. P-values were obtained using the non-parametric Mann-Whitney U test, and standard error bars are shown. A scatterplot of Tnc (B) and Tnfsf15 (C) expression versus Mcs5c copy number demonstrate no correlation between the two (Pearson correlation coefficient, R = -0.229 & 0.229, n = 10 & 9, p-value = 0.524 & 0.553, respectively). A linear trend line is shown with the dotted line (slope = -0.121 & 0.173, respectively).

(TIF)

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Author Contributions

Conceived and designed the experiments: ANH MNG.

Performed the experiments: ANH JDH.

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References

1. American Cancer Society. Cancer Facts & Figures 2015. Atlanta: American Cancer Society; 2015.

2. Biro FM and Deardorff J. Identifying opportunities for cancer prevention during preadolescence and adolescence: Puberty as a window of susceptibility. J Adolesc Health. 2013; 52: S15–S20. doi: 10.1016/j.jadohealth.2012.09.019 PMID: 23601607

3. Land CE, Tokunaga M, Koyama K, Soda M, Preston DL, Nishimori I, et al. Incidence of female breast cancer among atomic bomb survivors, Hiroshima and Nagasaki, 1950–1990. Radiat Res. 2003; 160: 707–717. PMID: 14640793

4. Hancock SL, Tucker MA, Hoppe RT. Breast cancer after treatment of Hodgkin’s disease. J Natl Cancer Inst. 1993; 85: 25–31. PMID: 8416252

5. Russo J, Wilgus G, Russo IH. Susceptibility of the mammary gland to carcinogenesis: I. Differentiation of the mammary gland as determinant of tumor incidence and type of lesion. Am J Pathol. 1979; 96: 721–736. PMID: 112872

6. Ariazi JL, Haag JD, Lindstrom MJ, Gould MN. Mammary glands of sexually immature rats are more susceptible than those of mature rats to the carcinogenic, lethal, and mutagenic effects of N-nitroso-N-methylurea. Mol Carcinogen. 2005; 46: 155–164.

7. Fachal L and Dunning AM. From candidate gene studies to GWAS and post-GWAS analyses in breast cancer. Curr Op Genet Dev. 2015; 30: 32–41. doi: 10.1016/j.gde.2015.01.004 PMID: 25727315

8. Russo J. Significance of rat mammary tumors for human risk assessment. Toxicol Pathol. 2015; 43: 145–170. doi: 10.1177/0192623314532036 PMID: 25714400

9. Samuelson DJ, Haag JD, Lan H, Monson DM, Shultz MA, Kolman BD, et al. Physical evidence of Mcs5, a QTL controlling mammary carcinoma susceptibility, in congenic rats. Carcinogenesis. 2003; 24: 1455–1460. PMID: 1284486

10. Samuelson DJ, Aperavich BA, Haag JD, Gould MN. Fine mapping reveals multiple loci and a possible epistatic interaction within the mammary carcinoma susceptibility quantitative trait locus, Mcs5. Cancer Res. 2005; 65: 9637–9642. PMID: 16266982

11. Veillet AL, Haag JD, Remfert JL, Meilahn AL, Samuelson DJ, Gould MN. Mcs5c: A mammary carcinoma susceptibility locus located in a gene desert that associates with Tenascin C expression. Can Prev Res (Phila). 2011; 4: 97–106.

12. Conover CA, Bale LK, Overgaard MT, Johnstone EW, Laursen UH, Fuchtbauer EM, et al. Metalloproteinase pregnancy-associated plasma protein A is a critical growth regulatory factor during fetal development. Development. 2004; 131: 1187–1194. PMID: 14973274

13. Monget P, Mazeron S, Delpuich T, Maurel MC, Manière S, Zapf J, et al. Pregnancy-associated plasma protein-A is involved in insulin-like growth factor binding protein-2 (IGFBP-2) proteolytic degradation in bovine and porcine preovulatory follicles: Identification of cleavage site and characterization of IGFBP-2 degradation. Biol Reprod. 2003; 68: 77–86. PMID: 12493698

14. Lawrence JB, Oxvig C, Overgaard MT, Sottrup-Jensen L, Gleich GJ, Hays LG, et al. The insulin-like growth factor (IGF)-dependent IGF binding protein-4 protease secreted by human fibroblasts is pregnancy-associated plasma protein-A. Proc Natl Acad Sci USA. 1999; 96: 3149–3153. PMID: 10077652

15. Laursen LS, Overgaard MT, See R, Boldt HB, Sottrup-Jensen L, Giudice LC, et al. Pregnancy-associated plasma protein-A (PAPP-A) cleaves insulin-like growth factor binding protein (IGFBP)-5 independent of IGF: Implications for the mechanism of IGFBP-4 proteolysis by PAPP-A. FASEB J. 2001; 15: 36–40. PMID: 11522292
16. Kleinberg DL, Wood TL, Furth PA, Lee AV. Growth hormone and Insulin-like growth factor-I in the transition from normal mammary development to preneoplastic mammary lesions. Endocr Rev. 2009; 30: 51–74. doi: 10.1210/er.2008-0022 PMID: 19075184

17. Ruan W and Kleinberg DL. Insulin-like growth factor I is essential for terminal end bud formation and ductal morphogenesis during mammary development. Endocrinology. 1999; 140: 5075–5081. PMID: 10537134

18. Bonnette SG and Hadsell DL. Targeted disruption of the IGF-I receptor gene decreases cellular proliferation in mammary terminal end buds. Endocrinology. 2001; 142: 4937–4945. PMID: 11606462

19. Christopoulos PF, Msaouel P, Koutsilieris M. The role of the insulin-like growth factor-1 system in breast cancer. Mol Cancer. 2015; 14: 10537134

20. de Ostrovich KK, Lambertz I, Colby JK, Tian J, Rundhaug JE, Johnston D, et al. Paracrine overexpression of insulin-like growth factor-1 enhances mammary tumorigenesis in vivo. Am J Pathol. 2008; 173: 824–834. doi: 10.2353/ajpath.2008.071005 PMID: 18688034

21. Boldt HB and Conover CA. Overexpression of Pregnancy-associated plasma protein-A in ovarian carcinoma cells promotes tumor growth in vivo. Endocrinology. 2011; 152: 1470–1478. doi: 10.1210/en.2010-1095 PMID: 21303951

22. Pan H, Hanada S, Zhao J, Mao L, Ma MZ. Protein secretion is required for pregnancy-associated plasma protein-A to promote lung cancer growth in vivo. PLoS One. 2012; 7: e48799. doi: 10.1371/journal.pone.0048799 PMID: 2352806

23. Ryan AJ, Napolitano S, Fitzpatrick PA, Currid CA, O’Sullivan NC, Harney JH. Expression of a protease-resistant insulin-like growth factor-binding protein 4 in human breast cancer. Br J Cancer. 2009; 100: 278–286. doi: 10.1038/sj.bjc.6605141 PMID: 19536088

24. Ciriello G, Gatza ML, Beck AH, Wilkerson MD, Rundhaug JE, Johnston D, et al. Comprehensive molecular portraits of invasive lobular breast cancer. Cell. 2015; 163: 506–519. doi: 10.1016/j.cell.2015.09.033 PMID: 26451490

25. Gao J, Aksoy BA, Dogrusoz U, Dresdner G, Gross B, Sumer SO, et al. Integrative analysis of complex cancer genomics and clinical profiles using the cBioPortal. Sci Signal. 2013; 6: p1t. doi: 10.1126/scisignal.2004088 PMID: 23550210

26. Cerami E, Gao J, Dogrusoz U, Gross BE, Sumer SO, Aksoy BA, et al. The cBio cancer genomics portal: An open platform for exploring multidimensional cancer genomics data. Cancer Discov. 2012; 2: 401–404. doi: 10.1158/2159-8290.CD-12-0095 PMID: 22588877

27. Irizarry RA, Ladd-Acosta C, Wen B, Wu Z, Montano C, Onyango P, et al. The human colon cancer methylome shows similar hypo- and hypermethylation at conserved tissue-specific CpG island shores. Nat Genet. 2009; 41: 178–186. doi: 10.1038/ng.298 PMID: 19151715

28. Lettice LA, Heaney SJ, Purdie LA, Li L, de Beer P, Oostra BA, et al. A long-range Shh enhancer regulates expression in the developing limb and fin and is associated with preaxial polydactyly. Hum Mol Genet. 2003; 12: 1725–1735. PMID: 12837695

29. Benko S, Gordon CT, Mallet D, Sreenivasan R, Thauvin-Robinet C, Brendehaug A, et al. Disruption of a long distance regulatory region upstream of SOX9 in isolated disorders of sex development. J Med Genet. 2011; 48: 825–830. doi: 10.1136/jmedgenet-2011-100255 PMID: 22051515

30. Smits BM, Haag JD, Rissman AI, Sharma D, Tran A, Schoenborn AA, et al. The gene desert mammary carcinoma susceptibility locus Mcs1a regulates Nr2f1 modifying mammary epithelial cell differentiation and proliferation. PLoS Genet. 2013; 9: e1003549. doi: 10.1371/journal.pgen.1003549 PMID: 23785296

31. Acevedo N, Reinius LE, Greco D, Gref A, Orsmark-Pietras C, Persson H, et al. Risk of childhood asthma is associated with CpG-site polymorphisms, regional DNA methylation and mRNA levels at the GSDMB/ORMDL3 locus. Hum Mol Genet. 2015; 24: 875–890. doi: 10.1038/hmg/ddu479 PMID: 25256354

32. Bockmühl Y, Patchev AV, Madejska A, Hoffmann A, Sousa JC, Sousa N, et al. Methylation at the CpG island shore region upregulates Ntrc1 promoter activity after early-life stress. Epigenetics. 2015; 10: 247–257. doi: 10.1080/15592294.2015.1017199 PMID: 25793778

33. Farkas SA, Böttiger AK, Isaksson HS, Finnell RH, Ren A, Nilsson TK. Epigenetic alterations in folate transport genes in placental tissue from fetuses with neural tube defects and in leukocytes from subjects with hyperhomocysteinemia. Epigenetics. 2013; 8: 303–316. doi: 10.4161/epi.23988 PMID: 23417011

34. Park JL, Kim HJ, Seo EH, Kwon OH, Lim B, Kim M, et al. Decrease of 5hmC in gastric cancers is associated with TET1 silencing due to with DNA methylation and bivalent histone marks at TET1 CpG island 3’-shore. Oncotarget. 2015; 6: 37647–37662. doi: 10.18632/oncotarget.6069 PMID: 26462176
35. Perisic T, Holsboer F, Rein T, Zschocke J. The CpG island shore of the GLT-1 gene acts as a methyla-
tion-sensitive enhancer. Glia. 2012; 60: 1345–1355. doi: 10.1002/glia.22353 PMID: 22593010
36. Rao X, Evans J, Chae H, Pilrose J, Kim S, Yan P, et al. CpG island shore methylation regulates caveo-
lin-1 expression in breast cancer. Oncogene. 2013; 32: 4519–4528. doi: 10.1038/onc.2012.474 PMID:
23128390
37. Sugawara H, Iwamoto K, Bundo M, Ueda J, Miyauchi T, Komori A, et al. Hypermethylation of serotonin
transporter gene in bipolar disorder detected by epigenome analysis of discordant monozygotic twins.
Transl Psychiatry. 2011; 1: e24. doi: 10.1038/tp.2011.26 PMID: 22832526
38. Yang R, Pfütze K, Zucknick M, Sutter C, Wappenschmidt B, Marime F, et al. DNA methylation array
analyses identified breast cancer-associated HYAL2 methylation in peripheral blood. Int J Cancer.
2015; 136: 1845–1855. doi: 10.1002/ijc.29205 PMID: 25213452
39. Wright JB, Brown SJ, and Cole MD. Upregulation of c-MYC in cis through a large chromatin loop linked
to a cancer risk-associated single-nucleotide polymorphism in colorectal cancer cells. Mol Cell Biol.
2010; 30: 1411–1420. doi: 10.1128/MCB.01384-09 PMID: 20065031
40. Palstra RJ, Tolhuis B, Splinter E, Nijmeijer R, Grosveld F, de Laat W. The β-globin nuclear compart-
ment in development and erythroid differentiation. Nat Genet. 2003; 35: 190–194. PMID: 14517543
41. Kleinberg DL and Barcellos-Hoff MH. The pivotal role of Insulin-like growth factor I in normal mammary
development. Endocrinol Metab Clin N Am. 2011; 40: 461–471.
42. Russo J, Tay LK, Ciocca DR, Russo IH. Molecular and cellular basis of the mammary gland susceptibi-
liity to carcinogenesis. Environmental Health Perspectives. 1983; 49: 185–199. PMID: 6403347
43. Interagency Breast Cancer and Environmental Research Coordinating Committee (IBCERCC). Breast
Cancer and the Environment: Prioritizing Prevention. 2013. Available: https://www.niehs.nih.gov/about/
assets/docs/breast_cancer_and_the_environment_prioritizing_prevention_508.pdf
44. Vijayendran M, Beach SRH, Plume JM, Brody GH, Philibert RA. Effects of genotype and child abuse on
DNA methylation and gene expression at the serotonin transporter. Front Psychiatry. 2012; 3:55. doi:
10.3389/fpsyt.2012.00055 PMID: 22707942
45. Voisin S, Almén MS, Moschonis G, Chrousos GP, Manios Y, Schiöth HB. Dietary fat quality impacts
genome-wide DNA methylation patterns in a cross-sectional study of Greek preadolescents. Eur J
Hum Genet. 2015; 23: 654–662. doi: 10.1038/ejhg.2014.139 PMID: 25074463
46. Hvid H, Ekstrøm CT, Vienberg S, Oleksiewicz MB, Klopfleisch R. Identification of stable and oestrus
cycle-independent housekeeping genes in the rat mammary gland and other tissues. Vet J. 2011; 190:
103–108. doi: 10.1016/j.vetj.2010.09.002 PMID: 20952231
47. Whitehead RH, Demmler K, Rockman SP, Watson NK. Clonogenic growth of epithelial cells from nor-
mal colonic mucosa from both mice and humans. Gastroenterology. 1999; 117: 858–865 PMID:
10500068
48. Smits BM, Traun BD, Devries TL, Tran A, Samuelson D, Haag JD, et al. An insulator loop resides
between the synthetically interacting elements of the human/rat conserved breast cancer susceptibility
locus MCS5A/Mcs5a. Nucleic Acids Res. 2012; 40: 132–147. doi: 10.1093/nar/gkr610 PMID:
21914726
49. Ran FA, Hsu PD, Wright J, Agarwala V, Scott DA, Zhang F. Genome engineering using the CRISPR-
Cas9 system. Nat Protoc. 2013; 8: 2281–2308. doi: 10.1038/nprot.2013.143 PMID: 24157548