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# NOVEL APPROACHES INVESTIGATING PREDISPOSITION FOR MATERNAL GROUP B STREPTOCOCCUS COLONIZATION

by

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> A Dissertation Submitted to the Graduate Faculty

> > of the

University of North Dakota

In partial fulfillment of the requirements

for the degree of

Doctor of Philosophy

Grand Forks, North Dakota May 2014

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This dissertation, submitted by Michelle Lynn Wright in partial fulfillment of the requirements for the Degree of Doctor of Philosophy from the University of North Dakota, has been read by the Faculty Advisory Committee under whom the work has been done, and is hereby approved.

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Michelle Lynn Wright February 15, 2014

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# ABSTRACT

Group B streptococcus (GBS) is the leading cause of infectious neonatal morbidity and mortality in the United States. Maternal GBS colonization is the primary risk factor associated with neonatal infection. However, maternal risk factors for GBS colonization are ambiguous. A conceptual framework of gene-environment interactions guided the approach for this study analyzing DNA methylation, serum cytokines, and vitamin D levels. The purpose of this study was to identify potential maternal biomarkers associated with GBS colonization. Descriptive statistics were conducted to depict sample characteristics (n=42 pregnant women) and identify potential confounding variables including, but not limited to: medical history, race, weight, and infections. A series of repeated measures ANOVAs were performed to compare each of three serum cytokines (TNF- $\alpha$ , IL-6 and IL-10) and vitamin D levels between the two groups in each trimester of pregnancy. All statistical analysis was completed using a two-tailed alpha of < 0.05 or 95% confidence interval. Mean differences of greater than 20% in DNA methylation of maternal white blood cells collected in the first trimester were analyzed using a false discovery rate of 0.05 to determine significance, as well as independent sample t-tests with a p-value of 0.05 using the Illumina Infinium platform and grouped by GBS status (n=9/group) identified in the third trimester. Function of differentially methylated genes was determined using DAVID Bioinformatics software to identify clinically relevant findings. No statistically significant differences in IL-6 F(2, 80) = 2.99, p = 0.056; IL-10 F(2, 80) = 0.445, p = 0.642; TNF- $\alpha$  F(2, 80) = 2.187, p = 0.119; or vitamin D F(1.380, p) = 0.1

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55.218) = 0.882, p = 0.384 were identified between GBS positive and negative women during pregnancy. Analysis of DNA methylation indicates there are no statistically significant differences between GBS positive and GBS negative women using and FDR of 0.05. When a less stringent p-value of 0.05 was applied, 125 CpG sites differed by 20% or more between GBS positive versus negative women and different results are yielded using multiple statistical approaches (GenomeStudio versus R). Functional analysis suggests genes with methylation differences in the cell morphogenesis cluster may be associated with GBS colonization, although the significance is questionable.

## CHAPTER I

## INTRODUCTION

Group B streptococcus (GBS) is the leading cause of neonatal morbidity and mortality due to infection in the United States (Phares et al., 2008). Maternal GBS colonization is the primary risk factor associated with the development of neonatal GBS sepsis (Verani, McGee, Schrag, & Division of Bacterial Diseases Centers for Disease Control and Prevention (CDC), 2010). While between 10 - 30 percent of pregnant women are colonized with GBS (Schrag et al., 2002), risk factors for maternal colonization are ambiguous and inconclusive in the literature. To prevent transmission of GBS, colonized women are normally given antibiotics during the intrapartum period which significantly reduces the incidence of early onset GBS infections in neonates (Verani et al., 2010). However, current screening techniques have a 10% false negative rate (Towers et al., 2010) and do not prevent preterm labor, miscarriages, and stillbirths caused by GBS colonization; nor do they reduce the incidence of late onset GBS sepsis in infants (Clifford, Garland, & Grimwood, 2011; Jordan et al., 2008; Verani et al., 2010). There is a compelling need to investigate genetic and environmental factors that may help identify biomarkers for colonization because GBS continues to cause poor pregnancy outcomes and is associated with the absence of definitive maternal risk factors for colonization. If genetic and environmental factors can be identified, early screening and effective interventions can be developed and implemented. Preliminary data from our laboratory indicated DNA methylation differences can be measured early in pregnancy

between women with and without late pregnancy GBS colonization. DNA methylation, an epigenetic modification that can result in altered gene expression and related protein production, has the potential to drastically impact health and alter disease susceptibility (Baccarelli, Rienstra, & Benjamin, 2010; Berger, Kouzarides, Shiekhattar, & Shilatifard, 2009; Rodenhiser & Mann, 2006). Differential DNA methylation in genes regulating immunity and inflammation could lead to varied levels of pro-inflammatory tumor necrosis factor-alpha (TNF- $\alpha$ ), interleukin-6 (IL-6), and anti-inflammatory interleukin-10 (IL-10). TNF  $\alpha$ , IL-6, and IL-10 are proteins that are produced in varying amounts in response to threats to the immune system and could be clinical laboratory indicators for GBS colonization (Berner, Welter, & Brandis, 2002; Fan et al., 2003; Madureira et al., 2011; Maisey, Doran, & Nizet, 2008; Mikamo, Johri, Paoletti, Madoff, & Onderdonk, 2004; Ng et al., 2003; Parameswaran & Patial, 2010; Puliti et al., 2002; Santhanam et al., 1991; Vieira et al., 1991). Additionally, serum markers associated with immune function and vitamin D (25[OH]D) status have previously been identified and utilized as prognostic indicators for infectious disease (Chesney, 2010; Fahey et al., 1990) and may be a cost effective clinical intervention if altered serum vitamin D (25[OH]D) levels are associated with GBS colonization status. The long term goal for investigating different exogenous and endogenous clinical indicators in women with and without GBS colonization is to identify factors that may help later identify a mechanistic explanation for maternal GBS colonization and to develop and implement targeted primary prevention strategies to reduce neonatal sepsis caused by GBS. The specific purpose of this study was to identify variants in maternal blood that are associated with maternal GBS colonization.

#### Significance

*Streptococcus agalactiae*, or group B β-hemolytic streptococcus, is a gram positive bacterium that causes a wide spectrum of illness in multiple clinical populations. In 2005, GBS was attributed as the cause of 21,500 infections and 1,700 deaths in the United States (Phares et al., 2008). However, the rates of GBS colonization, independent of infection, were not reported or required to be reported. Primarily a bacterium that only causes invasive disease in patients with altered immune function (Johri et al., 2006), GBS was first associated with neonatal sepsis and maternal infection in the 1970s and continues to be the leading cause of neonatal sepsis (Verani et al., 2010). Maternal colonization with GBS is the strongest predictor for the development of GBS sepsis in the neonate. Currently, there are no effective strategies for preventing maternal GBS colonization because of the inconsistent results obtained from epidemiological studies (Clifford et al., 2011; Kovavisarach, Ying, & Kanjanahareutai, 2007; Zusman, Baltimore, & Fonseca, 2006). Studies investigating the number of pregnancies, maternal age, race, ethnicity, and other maternal characteristic have failed to identify common maternal risk factors. The variability in potential predisposing factors has resulted in research efforts focused on preventing transmission and subsequent development of GBS infection in neonates.

Compounding the issue of GBS infection in neonates is the fact that their risk for infection continues into the first 3 months of life. GBS infections are categorized in neonates by time of disease onset after delivery as either early or late onset. Early onset neonatal GBS sepsis occurs within the first 7 days of life with onset usually occurring within the first 48 hours of life. The time period ascribed to late onset neonatal GBS

sepsis is an infection that develops after the first week of life up until the infant is 3 months of age (Verani et al., 2010). In an attempt to circumvent GBS transmission to the neonate from colonized women, clinical guidelines recommend maternal screening for GBS colonization via cultures obtained from recto-vaginal swabs between 35-37 weeks gestation. If maternal screening tests are positive for GBS colonization, the CDC recommends intravenous antibiotics administration after the start of labor and at least 4 hours prior to delivery to prevent transmission of GBS to the neonate (Verani et al., 2010). Implementation of the CDC guidelines has resulted in a 80% decrease in the incidence of early onset neonatal GBS sepsis, although rates of late onset GBS sepsis have been unaffected (Schrag & Verani, 2013). The continued prevalence of early and late onset neonatal GBS sepsis since the guideline implementation could be a result of the 10% false negative rate associated with maternal screening for GBS colonization (Towers et al., 2010). Of the infants who develop GBS infections, 61-82 percent of infants are born to mothers with negative GBS screening at 35-37 weeks gestation and 6.3% of infants are colonized by GBS despite administration of maternal antibiotic treatment (Lin et al., 2011). This may be in part because colonization with GBS can be transient, intermittent, or persistent. Maternal GBS status could change between the time of screening and delivery. Additionally, the infant may become infected from environmental exposure after delivery (Verani et al., 2010). The transient nature of GBS colonization suggests that environmental factors or immune response may play a critical role in maternal colonization, warranting further investigation into maternal risk factors and more reliable screening.

In order to reach the Healthy People 2020 goal of a 10% reduction in neonatal GBS infections (U. S. Department of Health and Human Services (HHS), 2013), alternative interventions are required for two primary reasons. First, the 10% false negative rate during screening needs to be reduced. Women who screen negative for GBS colonization will not receive antibiotic prophylactic treatment and could contribute to the continued prevalence of GBS disease in neonates. Furthermore, it is unclear why some women initially screen negative and later convert to GBS positive status. The transient nature of some GBS colonization implies there may be environmental factors that contribute to colonization susceptibility. Second, increasing rates of antimicrobial resistance to intrapartum antibiotics are being reported. The antibiotic currently recommended for treatment of GBS colonization during pregnancy is intravenous penicillin (Verani et al., 2010). However, approximately 10% of the general population report having a penicillin allergy (Solensky, 2003). Patients with allergies are administered erythromycin with clindamycin or vancomycin instead. In the US, over 54% of invasive strains of GBS are resistant to erythromycin and 33% are resistant to clindamycin (DiPersio & DiPersio, 2006) and similar rates of resistance are seen worldwide (Bergseng, Rygg, Bevanger, & Bergh, 2008; Janapatla, Ho, Yan, Wu, & Wu, 2008; Uh et al., 2007). Dual resistance to both clindamycin and erythromycin is also increasing, with rates reported as high as 94% of clindamycin resistant isolates also being resistant to erythromycin (Back, O'Grady, & Back, 2011). Additionally, Stoll et al. (2011) found 53% of infants that developed early onset sepsis were born to mothers who had received intrapartum antibiotic treatment. The failure rates associated with current treatment methods are associated with continuing neonatal morbidity and moratlity. New

strategies which are robust at identifying women at risk for GBS colonization may increase the likelihood of developing successful intervention alternatives, further reducing the incidence of neonatal GBS sepsis.

The significance of the problem surrounding maternal GBS colonization is compounded by the fact that current guidelines to prevent neonatal GBS do not prevent poor maternal and fetal outcomes associated with GBS colonization. Administering antibiotics during labor does not prevent stillbirths, miscarriages, chorioamnionitis, or other poor pregnancy outcomes associated with GBS prior to 35 weeks gestation. Furthermore, there has been a 32% increase in GBS infections in non-pregnant adults with no information on colonization rates available (Phares et al., 2008). It is unclear why there has been such a large increase in the incidence of GBS infections.

Ambiguous risk factors for colonization, increasing rates of bacterial resistance to antibiotics, increasing GBS infections in non-pregnant populations, and failure rates associated with current treatment methods are compelling reasons to discover new approaches to identify individuals at risk for GBS colonization. Identification of factors associated with GBS colonization will provide new clinically relevant targets to prevent and treat GBS colonization. Identification of definitive environmental and/or genetic maternal risk factors associated with GBS colonization is a substantively different approach to preventing neonatal GBS sepsis. This contribution is significant and will improve scientific knowledge by identifying key differences in pregnant women with and without GBS colonization. If differences are identified between women with and without GBS colonization, knowledge gained from this study can be used to develop a more

accurate screening tool or prevent maternal GBS colonization thereby improving clinical practice and pregnancy outcome.

#### Purpose

The purpose of this study was to identify variants in maternal blood that are associated with maternal GBS colonization by investigating the following specific aims:

- Differentiate serum levels of TNF-α, IL-6, and IL-10 as potential clinical laboratory indicators for GBS colonization longitudinally in pregnant women; and examine circulating 25-hydroxyvitamin D (25[OH]D) (vitamin D (25[OH]D) as a potential covariate of TNF-α, IL-6, and IL-10 serum levels.
- 2. Determine if DNA methylation are different in pregnant women with and without GBS colonization.
- 3. Examine the relationship of any differentially methylated genes for association with immune function and inflammatory serum markers in pregnant women colonized with GBS.

## **Conceptual Framework**

Investigations of the interaction between genomic and environmental factors have been suggested as key research avenues in identifying the most effective methods to prevent disease. Cohesive investigations illuminate biochemical explanations for health problems and identify modifiable risk factors that can be controlled or altered to prevent disease (Willett, 2002). The relationship between genetic and environmental factors associated with the development of disease was first suggested in 1902 (Hunter, 2005). Examples of conditions known to be affected by genetic and environmental factors include: halitosis (Bretz et al., 2011), head and neck cancer after human papilloma virus exposure (Jamaly et al., 2012), sun exposure and skin cancer (Rees, 2004), and an increased susceptibility to human immunodeficiency virus infection based on cytokine profiles (Smith et al., 1997). Hunter (2005) published a conceptual model (Figure 1)

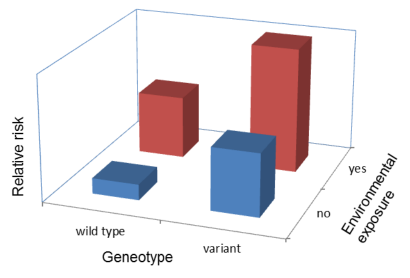


Figure 1. Gene-environment interaction.

illustrating how gene-environment interactions can potentiate disease processes. In the model, either a genetic variant or environmental exposure may result in disease even in the absence of an interaction between the two. The gene-environment model described by Hunter shows relative risk for disease states can be the result of either 1) a genetic variant 2) an environmental factor or 3) the interaction of the two. However, when there is an interaction between a predisposing genetic variant and environmental exposure, the risk for disease may be amplified. Hunter has used the inherited condition xeroderma pigmentosum to illustrate the model. Inheritance of the genetic mutation that causes the disorder greatly increases the risk for skin cancer and sun exposure further elevates risk (Cleaver, 2005; Kraemer, Lee, Andrews, & Lambert, 1994). When both elements are present, it results in a multiplicative effect drastically increasing disease risk (Figure 1). In other words the most basic interpretation of this relationship can be expressed as a

mathematical equation, genetic variants (G) x environmental exposure (E) = change in disease susceptibility (D):

#### $G \ge D$

Another example is that of emphysema. Individuals with the genetic mutation (G) causing an alpha 1-antitrypsin deficiency are likely to develop emphysema at a young age (DeMeo & Silverman, 2004). Smoking is an environmental exposure (E) linked to development of emphysema. An individual who has the mutation and smokes may develop more severe emphysema ( $\Delta D$ ) at an earlier age than a person with the mutation who does not smoke.

# Genetic Variants

The identification of the underlying genetic mechanism for an observable characteristic, referred to as a phenotype, is a common undertaking in the field of genetics. The genetic code contained within the nucleus of the cell is the genotype. The combination of dominant and recessive alleles, commonly called genes, and non-coding regions of the genome make up an individual's genotype. The genotype serves as a blueprint, contributing to the development of the phenotype. Copy number variants, insertions, deletions, and single nucleotide polymorphisms are some examples of how changes in the genetic code can lead to phenotypic changes. Hunter (2005) does not define genetic variants when presenting his model. However, the most simplistic model compares the "genotype" of individuals with or without a given trait to illustrate how environment may impact the observed phenotype based on genotype. The complexity of the model increases when multiple genes contribute to the development of a single phenotype. Eye color is an example of a complex phenotype resulting from multiple genes (Liu et al., 2009). Hunter (2005) did not include epigenetic modifications in his discussion of genetic variants, although he did acknowledge the plausibility of assessing factors altering gene expression in G x E interactions. Since the introduction of this simple model, the field has advanced to include methods for measuring epigenetic mechanisms that alter gene expression. Considering epigenetic alterations have the ability to augment phenotypic expression; inclusion in, or extension of, the model is logical.

## Environmental Variants

The environment has a profound impact on the health of individuals. While Hunter (2005) does not specifically define environmental variants in his model, he does consistently state that "environmental and lifestyle" factors must be assessed, implying a broad definition of environment. Ottman (1996) previously defined environmental exposures in gene-environment studies as:

The environmental risk factor can be an exposure, either physical (e.g., radiation, temperature), chemical (e.g., polycyclic aromatic hydrocarbons), or biological (e.g., a virus); a behavior pattern (e.g., late age at first pregnancy); or a "life event" (e.g., job loss, injury). This is not intended as an exhaustive taxonomy of risk factors, but indicates as broad a definition as possible of environmental exposures. (p. 764-765)

Further, epidemiologists are experts at identifying associations between environmental exposure and disease processes. Unfortunately, many epidemiological studies fail to collect DNA samples making it impossible to assess the genetic variation in large samples of unrelated subjects. Decreasing the existing disconnect between epidemiologic and genetic analysis is possible when factors identified in epidemiologic studies are

assessed in genetic research involving human subjects. Most genetic studies involving human subjects collect some information about participants. Data may be limited to demographic information or include detailed laboratory values and other potentially relevant risk factors. When designing genetic studies with human subjects, improved assessment and collection of environmental exposure data could offer insight into disease processes (Hunter, 2005).

### Gene-environment Interactions

Hunter (2005) describes two possible approaches for interpreting geneenvironment interactions that contribute to disease in the model. The interpretation of the interaction is dependent on the statistical model selected, and must be appropriate for the type of clinical question being addressed. When scientists are interested in determining how factors contribute to the relative risk of a disease, they assume a multiplicative interaction where the risk is either increased or decreased when multiple factors contribute to disease development. Results from studies using a multiplicative approach usually report findings in terms of relative risk. Using the emphysema example from before, the probability of developing emphysema early is more likely to occur if a person with the genetic defect smokes. The probability of how likely it is for an outcome to occur is referred to as the relative risk.

If the *a priori* assumption is the interaction is a joint effect, the relationship is additive. To illustrate this type of relationship a clinical example where diagnosis of a disease is usually based clinical presentation will be used as an exemplar. For example, a patient presents to their primary care provider with skin lesions, a fever, and headache and is subsequently diagnoses with varicella by the provider. Each symptom does not

does not cause varicella; it is caused by a virus that results in the presentation of a specific a set of symptoms. The collection of symptoms together results in the clinical presentation caused by the viral infection. Studies utilizing an additive approach methodology usually report findings as rate differences, such as 80% of patients with chicken pox have a headache. Hunter encourages explicitly stating if the relationship assumed for analysis is multiplicative (relative risk) or additive (percent affected) in order to appropriately replicate and compare research studies. Research investigating gene-environment interactions has the potential to improve and individualize patient care by improving understanding of disease susceptibility allowing for development of alternate treatment and prevention strategies.

#### Modified Conceptual Framework

Epigenomic research has led to a greater understanding of how our genes and environment contribute to complex disease processes. Epigenome adds the Greek prefix "epi" to genome and literally translates to above the genome (epi, n.d). The epigenome is fundamental for normal human development and contributes to what makes individuals unique. Epigenomics is the study of heritable alterations in the chromosomes which do not change the DNA sequence itself, but result in a specific phenotype (Berger et al., 2009). Structural and functional modifications of the epigenome modulate expression of the genes encoded by DNA. Nurses, as members of interdisciplinary teams, can use advances in epigenomic techniques to better assess levels of health and disease risk. Additionally, it is important when conceptualizing environment in the extended model that endogenous and exogenous environmental variants be considered. The environment should be considered anything outside of the DNA because any exposure to the DNA

could alter the epigenomic signature. Epigenomic alterations should be included in studies investigating gene-environment interactions due to the dynamic, and potentially reversible, nature of the epigenome that can be modulated by endogenous and exogenous influences throughout the lifespan (Feinberg, 2008).

Moffitt, Caspi, & Rutter (2006) argue against incorporating epigenetic mechanisms into gene-environment interaction models because the alterations modulate the effects of environment on gene expression and do not represent actual alterations in the genes or the DNA sequence. This is true in some cases, but not all. For example, women with breast cancer type 2 susceptibility protein (BRCA2) mutations have a higher risk for developing ovarian cancer (Kanchi et al., 2014; King, 2003; Welcsh, 2001). Additionally, hypomethylation of BRCA2 in ovarian tumor DNA has been associated with advanced tumor staging (Chan, Ozcelik, Cheung, Ngan, & Khoo, 2002). In other words, there is an existing genetic variant that results in increased risk of a disease and the severity of disease is modulated by epigenetic variants. However, during embryonic development tissue specific DNA methylation patterns are established across the genome (Hajkova et al., 2002) and the epigenetic signature can be altered by environmental exposures. Since epigenetic (eG) modifications can occur as a direct result of environmental exposures (E) resulting in altered susceptibility to disease ( $\Delta D$ ) without a genetic variant being present, the same conceptual model can be applied to disease states without mutations.

#### Summary of Framework

Since publication of Hunter's (2005) model, methods for quantifying epigenomic modifications that alter gene expression have emerged. Assessing the impact of

environment on epigenetic signatures without gene mutations can be expressed as:

$$eG \ge E = \Delta D$$

Since DNA methylation patterns can contribute to the development of complex disease states, epigenetic modifications appear to be a good fit for Hunter's model. The model is useful in guiding research studies analyzing DNA methylation patterns because the research can be initiated by identifying epigenetic and environmental components separately and then considering the interaction between them. When little is known about a disease process or risk factors for a disease are ambiguous, the investigator can initiate inquiry by looking for DNA methylation patterns that may be associated with the disease state. A limitation to this approach, as with any retrospective analysis, is that we cannot establish causation. However, effect size established from a small exploratory study can be used to estimate the sample size needed for a prospective study that incorporates environmental interactions.

Application of multiplicative gene-environment interaction models is useful in designing studies to determine environmental factors that modify epigenomic variants. DNA methylation, an epigenetic process that predominates during development and can be modulated throughout postnatal life, is influenced by factors such as nutrition, body weight and smoking status that are amenable to nursing interventions (Davis & Uthus, 2004; Kargul & Laurent, 2009). By modifying Hunter's (2005) model incorporating epigenome, instead of genotype, it is possible to identify epigenomic signatures, environmental factors, and interaction between the two that result in complex disease states.

The purpose of this study was to identify variants in maternal blood that are associated with maternal GBS colonization. In chapter two, what is known about GBS colonization is reviewed to justify the need for using a substantially different approach to identifying host factors that may be associated with GBS colonization. To date, epidemiological studies have failed to identify consistent maternal risk factors. The approach for this study will be to analyze epigenomic variants of affected individuals, specifically DNA methylation, and evaluate several maternal endogenous environmental variants IL-10, IL-6, TNF- $\alpha$ , and serum vitamin D (25[OH]D) (25[OH]D). Lastly, alterations in DNA methylation patterns will be evaluated to determine if they are involved in functional pathways associated with immune function. Identification of specific alterations in DNA methylation that result in altered immune function would suggest a mechanistic explanation as to why a third of pregnant women are colonized with GBS.

#### **Research Questions**

- Are serum levels of TNF-α, IL-6, IL-10, and vitamin D (25[OH]D) different in pregnant women with GBS colonization than pregnant women without GBS colonization?
- 2. Are DNA methylation patterns different between pregnant women with GBS colonization and those without GBS colonization?
- 3. Is there a relationship between differentially methylated genes and immune function in pregnant women colonized with GBS?

#### Assumptions

The study will be conducted based on the following pre-stated assumptions:

- Pregnant women presenting with GBS colonization did not have clinical symptoms indicative of infection such as fever, chorioamnionitis, bacturia or preterm labor at the time recto-vaginal swabs for culture were collected. Medical records indicate no other active infections at the time of screenings.
- Altered levels of circulating TNF-α, IL-6, IL-10, and vitamin D (25[OH]D) are associated with infectious disease susceptibility (Berner et al., 2002; Fan et al., 2003; P Madureira et al., 2011; Maisey et al., 2008; Mikamo et al., 2004; Ng et al., 2003; Parameswaran & Patial, 2010; Puliti et al., 2002; Santhanam et al., 1991; Vieira et al., 1991).
- Serum vitamin D (25[OH]D) levels are variable related to cytokine production (Hopkins et al., 2011; Shab-Bidar et al., 2012).
- Environmental influences throughout life have the potential to induce variation in DNA methylation patterns, modulating gene expression that contributes to health and disease states (Baccarelli et al., 2010; Berger et al., 2009; Rodenhiser & Mann, 2006)
- Differences in maternal DNA methylation patterns during early pregnancy among women with and without GBS colonization represent a biomarker for early risk identification.

#### CHAPTER II

### **REVIEW OF LITERATURE**

Maternal group B streptococcus (GBS) colonization continues to be a global health burden among pregnant women and neonates despite current treatment and prevention strategies because neither adequately address the underlying cause of disease (Edmond et al., 2012; Verani et al., 2010). In this chapter, evidence supporting the modification of the gene-environment framework, described in chapter 1, to incorporate the inclusion of epigenomic-environment interactions will be described for GBS colonization as the intended research target. The characteristics of GBS and clinical significance of GBS colonization and infection will also be reviewed. GBS continues to be a pathogen of interest due to increasing resistance to antibiotics, continued neonatal infections despite antibiotic prophylaxis, and increasing incidence of invasive disease in non-pregnant adults. Current treatment and prevention protocols have been in place since the 1990s and the incidence of disease has plateaued in the neonatal population, but has risen in historically unaffected populations (Edmond et al., 2012; Phares et al., 2008; Schrag & Verani, 2013). Most recently, multiple scientific teams have begun to develop vaccines against GBS in attempts to mitigate the disease burden caused by GBS (Johri et al., 2006; Schrag & Verani, 2013). However, characteristics of the bacterium and ethical issues have hindered successful vaccine development. The intent of this study was to identify variants in maternal blood that are associated with maternal GBS colonization

that could help identify new intervention targets to prevent GBS colonization and subsequent infection.

#### Group B Streptococcus

GBS is a gram positive bacterium that causes invasive diseases, such as pneumonia, meningitis, chorioamnionitis, and sepsis, primarily affecting pregnant women, the elderly, and infants. GBS continues to cause significant morbidity and mortality, particularly in neonatal populations, despite current clinical practice designed to prevent the transmission of GBS from mother to neonate during delivery (Phares et al., 2008). Clinical recommendations described in Chapter 1 to circumvent GBS transmission to the neonate, have significantly decreased the incidence of early neonatal sepsis. However, there have been no further decrease in the amount of late or early onset sepsis and new approaches for prevention infection are need to further reduce poor outcomes associated with GBS infection. What is known about GBS infection and colonization will be reviewed, as well as novel new approaches to identify endogenous maternal factors that may be associated with risk for GBS colonization.

#### Characteristics of GBS bacterium

GBS are facultative anaerobic gram positive cocci that grow in pairs or chains. Initially, GBS was only associated with cattle as the cause of bovine mastitis. Lancefield (1933), first categorized streptococci into five groups based on cell wall carbohydrate antigens, and the groups also differ in laboratory identification techniques, colony morphology, and disease association. The groups are labeled A-E, GBS belongs to Lancefield group B, producing 1-3 mm diameter grayish-white flat mucoid colonies when grown in the laboratory. The colonies have a narrow zone of hemolysis with a positive CAMP test. CAMP is an acronym for the scientist who developed the test (Christie, Atkinson, Munch, Peterson) for selectively identifying GBS. The additive they developed results in a larger area of clearance around the colonies because the additive increases the hemolytic activity of GBS (CDC, 2010). Group B specific antigen must also be detected when identifying GBS, usually done by latex agglutination. However, molecular identification via rapid polymerase chain reaction (PCR) and other methods is becoming more common (Bergseng et al., 2008; Kong, Gowan, Martin, James, & Gilbert, 2002).

# **GBS** Pathogenesis

The key feature of GBS that allows it to evade the host immune response is the thick capsular polysaccharide layer that surrounds the bacterium. Antigenic differences in the layer allows for differentiation into one of 10 distinct serotypes (Ia, Ib, II-IX) (Lancefield & Freimer, 1966; Slotved, Kong, Lambertsen, Sauer, & Gilbert, 2007). However, some GBS isolates are of indeterminate serotype which is likely because the antigens for the serotype have not yet been identified (Ferrieri, Baker, Hillier, & Flores, 2004). Prevalence of serotypes varies by geographic location (Ippolito et al., 2010), although serotype III causes most cases of disease in infants (48.9%) with serotype Ia (22.9%), serotype Ib (7.0%), serotype II (6.2%), and serotype V (9.1%) accounting for the majority of other cases (K M Edmond et al., 2012). The capsular polysaccharide layer prevents the binding of compliment factor C3 to the surface of GBS, allowing evasion of the immune recognition (Doran & Nizet, 2004; Spellerberg, 2000).

GBS beta-hemolysin/cytolysin ( $\beta$ -h/c) is the second virulence factor that is involved in GBS pathogenicity. GBS  $\beta$ -h/c is a non-immunogenic pore-forming

membrane associated toxin capable of damaging multiple tissues and impacts disease severity (Nizet et al., 1996; Puliti et al., 2000; Ring et al., 2002). In animal models GBS  $\beta$ -h/c resulted in increased bacterial load, pro-inflammatory cytokines IL-6 and IL-1 $\alpha$ , and mortality compared to non-hemolytic mutants (Puliti et al., 2000). However, when GBS  $\beta$ -h/c damages host cells it results in the release of IL-8 causing local inflammation and recruitment of neutrophils seen in GBS infections (Doran, Chang, Benoit, Eckmann, & Nizet, 2002). Additionally, macrophages exposed to GBS  $\beta$ -h/c have higher expression of nitric oxide synthase which generates 4 fold more nitric oxide than strains without  $\beta$ h/c. High levels of nitric oxide are exhibited in septic shock caused by GBS  $\beta$ -h/c (Ring et al., 2002).

The surface protein C5a peptidase plays a key role in adhesion to host cells and is present on all strains and serotypes of GBS (Cheng et al., 2001; G. Y.-H. Liu & Nizet, 2004). C5a peptidase is encoded by the ScpB gene and enables binding to epithelial cells (Brown et al., 2005). However, if the ScpB gene is deleted it does not completely inhibit GBS from adhering to host cells, suggesting that other factors play a role in GBS adherence to host cells (Cheng, Stafslien, Purushothaman, & Cleary, 2002; Lindahl, Stalhammer-Carlemalm, & Areschoug, 2005; Tamura, Hull, Oberg, & Castner, 2006). The following additional surface proteins also contribute to GBS adherence to host cells to varying degrees: pili,  $\alpha$ -C protein, Lmb, FbsA, and Rib. The surface proteins interact with fibronectin, fibrinogen, laninin, and integrins that attach to host cells (Doran & Nizet, 2004; Lindahl et al., 2005).

# Maternal GBS Colonization

Rates of maternal GBS colonization vary extensively worldwide (Table 1). Rates of GBS colonization have been reported as low as 1.8% in Maputo, Mozambique (de Steenwinkel et al., 2008), up to 65% in non-pregnant women in the United States (Meyn, Krohn, & Hillier, 2009). The gastrointestinal tract is thought to be the primary reservoir associated with maternal GBS colonization; sexual activity and abnormal vaginal microbiota presumably contribute to the development of vaginal GBS colonization (Meyn et al., 2009). GBS also is likely sexually transmitted because sexual partners are frequently colonized with the same strain (Foxman et al., 2006; Manning et al., 2004; Meyn et al., 2009; Meyn, Moore, Hillier, & Krohn, 2002). Maternal factors that have previously been associated with colonization are: young maternal age, black race, and having low levels of GBS-specific anticapsular antibodies (Verani et al., 2010). However, a study by Kovavisarach et al. (2007) identified older maternal age as a risk factor and Zusman et al. (2006) found no association with race or maternal age. Therefore, risk factors associated with maternal colonization based on epidemiological studies appear to differ by geographic location and are ambiguous (Kovavisarach et al., 2007; Phares et al., 2008; Verani et al., 2010; Zusman et al., 2006). Further research is necessary to identify factors associated with maternal GBS colonization so targeted prevention methods can be developed.

### GBS Disease in Pregnant Women

The incidence of invasive GBS disease in pregnant women is twofold higher than non-pregnant women (Deutscher et al., 2011). GBS can result in stillbirth, preterm birth, premature rupture of membranes, abortion, bacteremia, endometritis, chorioamnionitis,

Country	Colonization Rate (%)	Reference
Brazil	17.9	(Zusman et al., 2006)
Central African Republic	17.5	(Brochet, Couvé, Bercion, Sire, & Glaser, 2009)
Germany	16.0	(Brimil et al., 2006)
Korea	8.3	(Kim et al., 2011)
Lebanon	17.7	(Seoud et al., 2010)
Mozambique	1.8	(de Steenwinkel et al., 2008)
Netherlands	21.0	(Valkenburg-van den Berg et al., 2006)
New Zealand	22.0	(Grimwood et al., 2002)
Norway	34.8	(Hakon Bergseng, Bevanger, Rygg, & Bergh, 2007)
Senegal	20.0	(Brochet et al., 2009)
Switzerland	21.0	(Rausch, Gross, Droz, Bodmer, & Surbek, 2009)
Taiwan	6.2	(Yang et al., 2012)
Thailand	18.1	(Kovavisarach et al., 2007)
United Kingdom	21.3	(N. Jones, Oliver, Jones, Haines, & Crook, 2006)
United States	24.2	(Verani et al., 2010)
Uruguay	17.3	(Laufer et al., 2009)
Zimbabwe	31.6	(Moyo, Mudzori, Tswana, & Maeland, 2000)

# Table 1. Global Maternal GBS Colonization Rates

pneumonia, puerperal sepsis, endocarditis, and infections of the genital tract, placenta, and amniotic sac. There is currently no known way to prevent GBS colonization and GBS screening is not conducted until late in pregnancy. Identification of maternal factors associated with risk for colonization may allow for the development of prevention and treatment strategies to prevent poor early pregnancy outcomes associated with GBS colonization. New prevention and treatment strategies are necessary because 70% of women who have poor pregnancy outcomes due to GBS will also endure poor fetal outcomes for their offspring (Phares et al., 2008; Verani et al., 2010).

# Maternal GBS Screening and Treatment

The CDC currently recommends screening for GBS colonization 35-37 weeks into the pregnancy for all pregnant women. Women positive for GBS are treated with antibiotics after they go into labor, preferably at least 4 hours prior to delivering the infant. Penicillin G (5 million units) and ampicillin (2 grams) are first line antibiotics used to prevent neonatal sepsis in infants born to mothers with GBS colonization because they both reach minimum bacteriocidal concentrations in the amniotic fluid, maternal, and fetal circulations (Pacifici, 2006; Verani et al., 2010). The following medications are recommended for GBS prevention for women with severe penicillin allergy: cefazolin, clindamycin, erthyromycin, and vancomycin. However, drug levels of these antibiotics are lower in fetal serum than maternal serum or have variable transfer rates across the placenta (Pacifici, 2006; Philipson, Sabath, & Charles, 1973). Ampicillin administered intravenously exceeds the minimum bactericidal concentration to kill 99.9% of GBS within five minutes of intravenous administration in the maternal and fetal circulation (Bloom, Cox, Bawdon, & Gilstrap, 1996). Penicillin levels 179 times above the minimum concentration required to eliminate GBS have been collected in fetal serum (Barber, Zhao, Buhimschi, & Illuzzi, 2008), indicating intrapartum antibiotics result in significant

maternal and fetal exposure from the intrapartum antibiotics administered. Neonatal serum levels of ampicillin are higher than maternal concentrations after delivery and persist for at least 5.6 hours (Colombo, Lew, Pedersen, Johnson, & Fan-Havard, 2006). Further studies evaluating the persistence of antibiotics in the fetal circulation after delivery are sparse, but likely contribute to the decreased incidence in early onset neonatal GBS sepsis.

*Maternal Vaccination:* Currently, there are a number of clinical trials underway examining proposed GBS vaccines for immunization of women prior to, or during, pregnancy. The vaccines currently being tested target either GBS capsular carbohydrates or proteins (Heath, 2011; Johri et al., 2006). Immunization with the capsular carbohydrate alone proved not to be sufficiently immunogenic. However, when capsular carbohydrate is combined with tetanus toxoid conjugate vaccine sufficient antibodies are produced against GBS (Baker, Rench, & McInnes, 2003). Baker at al.'s (2003) study only added conjugated type III capsular carbohydrate to the tetanus toxoid and did not result in immunity to other GBS serotypes. Ongoing studies are investigating potential capsular carbohydrate vaccines that are multivalent to ensure broader coverage to prevent infection. Despite the success of generating some immunity to GBS, a number of concerns related to GBS vaccine development remain. For example, vaccines may not be effective globally since there is variation in risk factors in the literature. There are also significant ethical concerns related to testing vaccines on pregnant women that could harm to the fetus (Johri et al., 2006; Paradiso, 2001). Vaccine manufacturing companies directly contribute to a lag in vaccine development for pregnant women because of fears of liability if the exposed child develops health issues later in life (Kaposy & Lafferty,

2012). Liability fears related to administering vaccines during pregnancy may worsen in light of growing evidence illustrating altered epigenomic patterns resulting from chemical exposures occurring *in utero*. For example, permanent alteration in methylation patterns can occur in fetal DNA in response to chemical exposures *in utero*, such as DES described previously. Litigation has also resulted from residual DES effects on the grandchildren of women given the medication during pregnancy (Rothstein, Cai, & Marchant, 2009). In light of this, vaccines and systemic antibiotic treatment to prevent GBS transmission may not be the least harmful approach. Further research is needed to determine the long term impact of vaccine and antibiotic administration in utero and early in development.

### GBS Disease in Neonates

GBS remains the leading cause of neonatal infectious morbidity and mortality, despite the administration of antibiotics colonized women to prevent vertical transmission (Clifford et al., 2011; Verani et al., 2010). The incidence of GBS disease in neonates is lower in developed countries, 0.4 - 0.81 per 100 live births, than in developing countries 0.91 - 1.81 per 100 live births (Table 2). Neonatal fatality rates are also disproportionately higher in some developing nations (Table 2), which could be due to variable prevalence of serotypes by geographic location (Johri et al., 2006) and different standards of medical care (Edmond et al., 2012; Heath, 2011).

Meningitis, bacteremia, and pneumonia are the neonatal clinical diagnoses most commonly caused by invasive GBS. In the US, the incidence of invasive neonatal GBS infection is higher in African American infants (Phares et al., 2008). A meta-analysis completed by Edmond et al. (2012) indicated that infants weighing less than 1.5 kg at

Country	Incidence/1000	Fatality Rate (%)	Reference
Denmark	0.4	8.0	(Ekelund & Konradsen, 2004)
Jamaica	0.91	3.6	(Trotman & Bell, 2006)
Malawi	1.81	33.0	(Gray, Bennett, French, Phiri, & Graham, 2007)
Netherlands	0.56	12.3	(Trijbels-Smeulders et al., 2007)
Norway	0.66	6.5 (1996-2005) 20.0 (2006)	(Bergseng et al., 2008)
United Kingdom and Ireland	0.72	9.7	(Heath et al., 2004)
United States	0.81-0.68	5.0-9.0	(Phares et al., 2008)

Table 2. Global Neonatal GBS Morbidity and Mortality

birth are 8 times more likely to develop invasive illness caused by GBS than normal weight infants. Additionally infants between 1.4 - 2.5 kg at birth are three times more likely to develop GBS infection. Neonatal GBS disease is classified by time of invasive disease onset after birth. Early onset GBS disease occurs during the first 7 days of life. Late onset GBS disease occur after the first week of life through the first 90 days after birth (Verani et al., 2010). In the US, the number of cases of early onset (1232) and late onset disease (1036) are nearly equal since the CDC guidelines for intrapartum antibiotics were implemented in the 1990s. However, mortality is higher for infants with early onset GBS disease (83 versus 48 deaths) (Phares et al., 2008).

### Early Onset Neonatal GBS Disease

The incidence of early onset GBS disease in neonates has decreased from 1.7 cases per 1,000 live births in 1990 to 0.37 cases per 1,000 live births in 2008 in the US; as a result of widespread implementation of intrapartum antibiotic administration for mothers colonized with GBS. Maternal GBS colonization is the strongest predictor of early onset disease. Other risk factors for early onset neonatal disease are GBS in maternal urine at any point in pregnancy, rupture of membranes greater than 12 hours, delivery before 37 weeks gestation, young maternal age, African American race, infection, low maternal anticapsular antibodies to GBS, prior delivery with GBS, and maternal fever greater than 37.5°C during labor (Verani et al., 2010). Women with heavy GBS colonization are more likely to infect their infants versus women who have a lower bacterial load (Regan et al., 1996; Yancey, Duff, Kubilis, Clark, & Frentzen, 1996). In other words, women with more GBS present have a higher likelihood of infecting their infants with GBS. A majority of infants that develop early onset disease are full term (77%) and 90% of infants become ill within 12 hours of birth (Phares et al., 2008). The most common presentation of early onset disease are pneumonia and sepsis, and less commonly meningitis (Verani et al., 2010).

### Late Onset Neonatal GBS Disease

The incidence of late onset GBS disease in neonates is currently 0.35 cases per 1,000 live births (Jordan et al., 2008). Intrapartum antibiotic administration for GBS colonization has not had any significant effect on the incidence of late onset disease (Berardi et al., 2013; Phares et al., 2008). Late onset disease commonly presents as bacteremia and meningitis, and less commonly pneumonia or local site infections such as

cellulitis (Berardi et al., 2013; Jordan et al., 2008). African American infants are disproportionately affected (Jordan et al., 2008). Unlike the presentation and time of onset of early onset disease, late onset disease is different between term and preterm infants; suggesting that there may be different mechanisms involved. For example, Berardi et al (2013) identified term infants present with late onset disease earlier than preterm infants. Additionally, the etiology may be different from early onset disease because less than 30% of infants who develop late onset disease had mothers with positive GBS screenings. It is been speculated the source may be breast milk, persistent maternal colonization (for the 30% where maternal GBS colonization was present), or healthcare workers but additional studies are needed for validation. Further, Jordan et al. (2013) identified that 47% of infants that developed late onset disease had been exposed to intrapartum antibiotics for either GBS colonization, Cesarean section, or other complications. Late onset disease has not been studied with the same intensity as early onset disease. Now that the incidence of early and late onset neonatal disease is equivalent, the depth of research investigating factors associated with late onset neonatal GBS may improve.

### GBS Infection in Non-pregnant Populations

The incidence of GBS infections in non-pregnant adults doubled from 3.6 per 100,000 people in 1990 to 7.3 per 100,000 people in 2007 (Skoff et al., 2009). Traditionally, GBS disease was seen in individuals with compromised immune systems due to advanced age or other underlying conditions. Similar to neonatal GBS disease, bacteremia is a common outcome of invasive GBS in adults followed by skin and/or soft tissue infections, and pneumonia. Since the implementation of intrapartum antibiotics for GBS colonized mothers, half of case fatalities are adults older than 65 years old (13.2% of those infected perish) and are approaching pneumonia fatality rates (20.6%) for the elderly population (Edwards, Rench, Palazzi, & Baker, 2005; Schrag et al., 2000). Colonization rates are similar to those seen in pregnant women (21.7%) and almost half of the elderly (47.3%) affected are colonized with serotype V (Edwards et al., 2005), whereas over 60% of disease in neonates is caused by serotype III and Ia (K M Edmond et al., 2012). Since this incidence is rising in this population, identification of factors contributing to maternal GBS colonization may also be informative for reducing the incidence in the elderly population as well.

### GBS Resistance to Antibiotics

As previously described, infants born to mothers adequately treated with intrapartum antibiotics still develop GBS disease and a large proportion of infants are born to mother with negative GBS screening. Additionally, GBS positive women treated with intrapartum antibiotics are more than 4 times likely to be positive for GBS 6 weeks after delivery than women not given intrapartum antibiotics (Manning et al., 2008). Interestingly, 65% of women in Manning et al.'s (2008) study continue to have GBS colonization despite antibiotic treatment and 18.3% of women that retained GBS positive status were colonized with a different strain of GBS. Perhaps the persistence of GBS colonization in mothers despite antibiotic treatment is a contributing factor in the unchanged incidence in late onset neonatal GBS disease since the implementation of intrapartum antibiotic protocols. Conversely, GBS bacteria may just be resistant to the antibiotic treatment. In recent years, there has been a push in implementing antibiotic stewardship programs in hospital settings to decrease the rates of antibiotic resistance. The programs should also incorporate GBS prophylaxis recommendations and providers should be aware of the resistance rates in their area. Many studies evaluating antibiotic resistance of GBS assert the strains remain susceptible to penicillin and amoxicillin (Castor et al., 2008; Chohan, Hollier, Bishop, & Kilpatrick, 2006; Garland et al., 2011). However, reduced susceptibility to penicillin by GBS has been identified in Hong Kong and warrants further monitoring for increasing resistance (Chu et al., 2007). Antibiotic sensitivity testing is particularly warranted for women with penicillin allergies, since clindamycin and erythromycin resistance is high (Table 3). Notably in the US, strains that are clindamycin or erythromycin resistance are likely to have dual resistance to both erythromycin (94.3%) and clindamycin (71.5%), respectively. In light of increasing

Country	Ε	С	EC	Reference	
Percent (%) Resistant					
Australia	17	22	38	(Garland et al., 2011)	
Korea	9.7	6.8	-	(Uh et al., 2007)	
Malawi	21	-	-	(Gray et al., 2007)	
New Zealand	-	15.4	7.7	(Grimwood et al., 2002)	
Norway*	11.9	10.9	25.4	(Bergseng et al., 2008)	
Taiwan	44.0	39.0	-	(Janapatla et al., 2008)	
United States	54.0	33.0	-	(DiPersio & DiPersio, 2006)	
United States (New York)	50.7	38.4	94.3/71.5	(Back et al., 2011)	

**Table 3. Global GBS Resistance to Antibiotics** 

E = Erythromycin

C = Clindamycin

EC = Both Clindamycin and Erythromycin

\*= E & C reported for adult cases EC reported for neonatal disease

antibiotic resistance, identification of definitive environmental and/or genetic maternal risk factors associated with GBS colonization could result in alternate clinical approaches for preventing neonatal GBS sepsis.

Epigenome-Environment Interaction and GBS Colonization Epidemiologic findings suggest an association between environmental factors and epigenetic alterations, serving as the basis for many complex diseases including obesity, type 2 diabetes, asthma, autism spectrum disorder, and attention deficit hyperactivity disorder (Latham, Sapienza, & Engel, 2012). Environmental influences that range from behavioral conditions, dietary factors, and toxic exposures can be modulated to achieve optimal health outcomes for the populations. Additionally, infections and events occurring within a person can alter epigenetic signatures (Tolg et al., 2011). Because of this, measuring endogenous factors that may contribute to disease development are also valuable in epigenomic-environment interactions studies. Further, if modifiable exogenous environmental factors can be linked to an altered endogenous environment or epigenome, implications for treatment and prevention are ascertained.

Epidemiologists have identified associations between environmental exposures and disease for decades linking poor hand hygiene of healthcare workers with patient sepsis in the 19th century (Gould, 2010), poor nutrition during pregnancy to cardiovascular disease later in life in offspring (Barker & Osmond, 1986; Barker, 1995), and smoking to lung cancer (Doll & Hill, 1950). However, epidemiological studies traditionally do not identify biologic mechanisms that cause disease and are limited to identifying the association between population-level risk factors and disease. Determining the role of DNA methylation in the origin of disease could help identify

epigenetic linkages, not explained by a change in sequence or genetic alteration, that have been elusive for most complex diseases. Factors identified in epidemiologic studies combined with DNA methylation analyses have the potential to identify epigenomicenvironment interactions at critical time points across the lifespan that contribute to disease phenotype and inheritance. Analysis of DNA methylation patterns alone will not explain why or how phenotypes are altered; nor will they identify effective primary, secondary or tertiary intervention strategies. Environmental factors (e.g., diet, lifestyle, stress) must be measured to identify interactions that may cause alterations in DNA methylation. Clinical assessments completed by nurses, particularly of lifestyle factors that alleviate or exacerbate symptoms, can help identify modifiable factors that are related to altered DNA methylation patterns. Each person's DNA and environmental exposures are unique and assessing both simultaneously will result in more personalized healthcare. For example, clinical assessments first recognized the association between diethylstilbestrol (DES) administration to prevent miscarriage and development of reproductive tract anomalies in offspring. *In vitro* and in adults, DES did not cause alteration in methylation patterns. However, DES causes hypermethylation of a gene responsible for reproductive tract development to organisms exposed *in utero* that persisted into adulthood (Bromer, Wu, Zhou, & Taylor, 2009). Therefore, it is necessary to perform detailed assessments of patient, and family, medical history to identify how exposures may alter gene expression because the associations are not always obvious.

To illustrate the utility of evaluating epigenomic changes in the context of geneenvironment interaction models as described in Chapter 1, the concept of DNA methylation and how perturbations in DNA methylation patterns can alter gene

expression and disease susceptibility will be reviewed. DNA methylation is the best understood epigenetic mechanism that modulates gene expression (Baccarelli, Rienstra, & Benjamin, 2010). Differential methylation induced by endogenous or exogenous influences can lead to both genome instability and inappropriate gene transcription, contributing to pathology. DNA methylation patterns are often specific to lineage, organ, and cell-type (Cedar & Bergman, 2012). For example, all the human cells in an individual contain the exact same DNA. The epigenomic signature of a cell will program it to differentiate into a heart, eye, or skin cell. Abnormal loss or gain of methylation at key DNA sites may result in inappropriate expression of a gene. When specific patterns of methylation are associated with a phenotype, such as risk for disease, the patterns can be used to identify those at risk for poor health outcomes and provide the basis for new treatments. The investigation of epigenetic markers to identify biological mechanisms of complex disease processes, such as atherosclerosis(Guay et al., 2013; Lund et al., 2004; Zaina, Lindholm, & Lund, 2005), schizophrenia(Auta et al., 2013; Costa et al., 2002; Sharma, 2005), and lupus(Absher et al., 2013; Li, Gorelik, Strickland, & Richardson, 2014; Sekigawa et al., 2003), have been increasing in recent years (Bergman & Cedar, 2013; Petronis, 2010; Rakyan, Down, Balding, & Beck, 2011; Rodenhiser & Mann, 2006). Furthermore, DNA methylation is heritable during cellular reproduction, and likely from one generation to another. This means methylation signature can pass from both cell to cell and transfer from parent to offspring (Guerrero-Bosagna & Skinner, 2012; Rodenhiser & Mann, 2006). Therefore, DNA methylation patterns passed across generations may provide an explanation for the transmission of complex disease susceptibility among families that is modulated by environmental exposures (Figure 2).

Figure 2 illustrates how alterations in DNA methylation by exogenous and endogenous factors can contribute to level of susceptibility to complex disease in individuals, with transmission of the pattern throughout future generations (Wright, Ralph, Ohm, & Anderson, 2013).

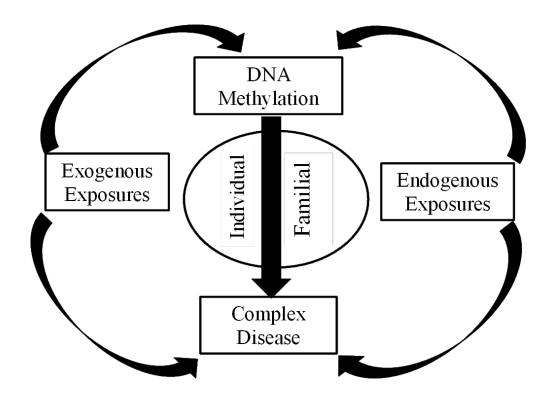


Figure 2. Influence of DNA Methylation in Complex Disease States.

DNA methylation patterns associated with GBS colonization could not be identified in the literature. However, altered DNA methylation patterns are known modulators of immune function and alterations have been associated with other infectious disease processes (Table 4). Notably, T-lymphocyte cell function and cytokine expression are altered by methylation patterns present on T cells (Fitzpatrick, Shirley, & Kelso, 1999). Additionally, DNA methylation patterns are altered by bacterial infection (Mikovits et al., 1998; Tolg et al., 2011) and different methylation patterns associated with the development of disease have been identified in animal models when they are colonized with altered bacterial populations after birth (Olszak et al., 2012). Since DNA methylation vitally contributes to cell differentiation in the immune system and programming memory in immune cells, it is an excellent candidate for identifying unknown mechanisms that may be associated with infectious diseases susceptibilities.

Immune Function or Alteration	Reference
Bacterial infection induces hypermethylation	(Tolg et al., 2011)
Discriminates between regulatory and conventional T cells	(Baron et al., 2007)
Downregulation of IFN- $\gamma$ in fetus, helps prevent fetal loss	(White, Watt, Holt, & Holt, 2002)
IgE production	(Liu, Ballaney, Al- alem, & Quan, 2008)
Maintenance of T cell memory and cytokine expression pattern	(Fitzpatrick et al., 1999)
Maternal bacterial infection promotes fetal hypermethylation	(Bobetsis et al., 2007)
Number and function of regulatory T cells	(Schaub et al., 2009)
Viral infection increases DNA methylation	(Mikovits et al., 1998)

#### **Table 4. DNA Methylation and Immune Function**

### DNA Methylation

DNA methylation is an epigenetic modification that may result in gene silencing, gene activation resulting in chromosome instability, inappropriate gene expression, and inability to carry the epigenomic signature to future cell lines (Bergman & Cedar, 2013; Cedar & Bergman, 2012; Jones, 2012; Rodenhiser & Mann, 2006). Epigenetic changes are heritable alterations in the chromosome that do not change the DNA sequence that result in a specific phenotype, which are observable characteristics (Berger et al., 2009;

He, Chen, & Zhu, 2011; Tost, 2010). DNA methylation is a specific epigenetic alteration in which a methyl group attaches to a cytosine (C) residue in DNA that is followed by a guanine (G) residue connected by a phosphate bond, commonly referred to as a CpG dinucleotide, and is currently the most well understood epigenetic mechanism (Baccarelli et al., 2010; Cedar & Bergman, 2012; Chen & Riggs, 2011; P. A. Jones, 2012). Areas with dense concentrations of CpG dinucleotides are located in promoter regions of genes, which is where transcription factors bind to initiate the reading of a DNA sequence so a gene will be expressed. Methylation of cytosines located in the promoter region can alter gene expression by blocking transcription of the DNA, resulting in gene silencing. Conversely, a loss of methylation at these sites may result in inappropriate expression of a gene.

Between 60-90 percent of cytosines are methylated in human DNA (Ehrlich et al., 1982) and the establishment of "normal" DNA methylation patterns are necessary for embryonic development. Normal patterns of methylation are required for the differentiation of cell types. For example, every cell in an individual's body has the same DNA sequence and methylation patterns present on the DNA sequence is specific to tissue type(Cedar & Bergman, 2012; Jones, 2012; Laird, 2010). In other words, the methylation pattern for a cell in the heart will be different from the methylation pattern on a cell in the eye, even though the DNA sequence is exactly the same. Alterations in methylation patterns also explain some of the processes that occur in complex disease states, such as delayed onset disease or situations where only one identical twin develops cancer (Boks et al., 2009; Fraga et al., 2005; Kaminsky et al., 2009; Petronis, 2001).

environmental exposures (Table 5), implying that we have the ability to intervene to prevent disease or promote desired health outcomes. Patterns of methylation associated

Environmental factor	Reference
Alcohol	(Choi et al., 1999)
Bisphenol-A (BPA)	(Bromer, Zhou, Taylor, Doherty, & Taylor, 2010)
Diethylstilbestrol (DES)	(Bromer, Wu, Zhou, & Taylor, 2009)
Exercise	(Barrès et al., 2012)
Fear	(Miller & Sweatt, 2007)
Hydralazine	(Cornacchia et al., 1988)
Maternal Care	(Weaver et al., 2004)
Maternal Diet	(Wolff, Kodell, Moore, & Cooney, 1998)
Microbiome	(Olszak et al., 2012)
Procainamide	(Cornacchia et al., 1988b)
Smoking	(Toyooka et al., 2003)
Traffic pollution	(Baccarelli et al., 2009)

 Table 5. Environmental Exposures that Alter DNA Methylation

with a specific phenotype, such as susceptibility to infection, have potential for use in identifying people at risk for developing conditions and as treatment targets. Leukemia treatments based on epigenetic markers have been approved and used successfully in the clinical setting (Rodriguez-Paredes & Esteller, 2011). Identifying differences in DNA methylation patterns during early pregnancy in women with GBS colonization could represent a biomarker for early risk identification or develop methods to prevent colonization. Furthermore, because there is a strong relationship between DNA

methylation patterns and the function of cells in the immune system (Table 4); differential DNA methylation in genes associated with immune function could offer mechanistic insight as to why certain bacteria, like GBS, colonize some individuals and not others.

### Environmentally Induced Modification

Environmental influences throughout life have the potential to induce variation in DNA methylation patterns, modulating gene expression that contributes to health and disease states (Baccarelli et al., 2010; Berger et al., 2009; Rodenhiser & Mann, 2006). For example, permanent alteration in methylation patterns can occur in fetal DNA in response to chemical exposures *in utero*. Diethylstilbestrol (DES) is a synthetic estrogen that was administered to pregnant women to prevent spontaneous abortions prior to the mid 1970's. DES causes hypermethylation of homeobox protein Hox-A10 (HOXA10), a gene that controls uterine organ development, resulting in reproductive tract anomalies that persist into adulthood (Bromer et al., 2009). Furthermore Bromer et al., determined the hypermethylation of HOXA10 was specific to the fetus and did not occur laboratory experiments using cell line or the pregnant women who received DES.

Assessment of DNA methylation patterns in disease states where the mechanism that alters gene expression are unknown can help identify etiology of disease. For example, in 1915, Kendall determined the microbes that colonize the gastrointestinal tract at birth are involved in normal development of the immune system. However, the biological mechanism of how this occurred remained unknown for years. Olszak et al. (2012) recently published a study suggesting that the type of bacteria that colonize the gastrointestinal tract in the neonatal period has an effect on the function of cells in the

immune system. This study was performed on germ-free and specific-pathogen free mice. Further, Olszak's study suggests that microbial exposure alters gene expression in specific tissues. The authors of the study noted hypermethylation of CpG sites in colonand lung tissues of the chemokine (C-X-C motif) ligand 16 (CXCL16) gene occurred when specific pathogens were not present during development. The CXCL16 gene encodes a chemokine receptor on invariant natural killer T cells (iNKT), resulting in higher accumulations of iNKT cells that are involved with inflammatory processes. Additionally, the higher accumulation of the iNKTs only occurred in the colon and lung when specific bacteria were not present. The authors hypothesize an environmental exposure later in life triggers various inflammatory disease processes programmed by the methylation changes in the bowel and lungs, like asthma and irritable bowel syndrome. Therefore, exposure to bacteria early in development affects the programming of the immune system, in mice, by causing perturbation in DNA methylation patterns. They concluded the findings could be extrapolated to humans because the mouse model used is similar to human cells. Further studies investigating alterations of methylation patterns in humans may be needed to demonstrate the effects of environmental exposures on immune function. This study aims to identify differential DNA methylation patterns in pregnant women colonized with GBS. If differential patterns are identified, future investigations will be focused on identifying causes of the altered methylation patterns.

### Endogenous Maternal Environment

Other clinical indicators could potentially be used to identify pregnant women at increased risk for GBS colonization and have not been discussed in published literature. Clinical indicators, such as serum cytokine levels that reflect immune system functioning, could also be associated with altered DNA methylation patterns. Variability in serum markers associated with immune function and vitamin D (25[OH]D) status have previously been identified and utilized as prognostic indicators of disease states, such as respiratory infections (Chesney, 2010), human immunodeficiency virus infections (Fahey et al., 1990), pancreatitis (Pezzilli et al., 1995), and depression (Kiecolt-Glaser & Glaser, 2002). There has been a plethora of research investigating the role of vitamin D (25[OH]D) as an immune function modulator in recent years (Figure 3) and could offer a cost effective intervention target if low serum vitamin D (25[OH]D) levels are associated with GBS colonization . In this section, a brief review of immune function during pregnancy, relevant cytokines, and vitamin D (25[OH]D) will be discussed.

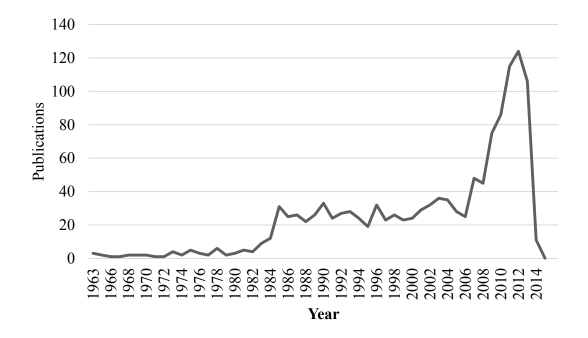


Figure 3. PubMed results for "Vitamin D" and "Immune Function". *Immune System During Pregnancy* 

Multiple alterations in immune function are necessary during pregnancy to prevent the mother's body from recognizing the developing fetus as a foreign pathogen.

Serum markers of immune function are increasingly being evaluated in pregnancy because alterations occur throughout normal pregnancy and inappropriate levels contribute to the development of pathology during pregnancy (Ponsonby, Lucas, Lewis, & Halliday, 2010). Alterations in the maternal immune system vary during pregnancy and can be effected by the overall health of the mother. For example, if a woman has a preexisting autoimmune disorder, like rheumatoid arthritis, she may experience remission from symptoms during the pregnancy as a result of altered immune functioning that protects the fetus. However, pregnancy does not result in the same altered state throughout the entire pregnancy because there are three distinct phases. First, in the early stages of pregnancy the environment is that of an invasion (Ashkar, Di Santo, & Croy, 2000; Dekel, Gnainsky, Granot, & Mor, 2010; Shimada et al., 2006). Implantation occurs, the placenta develops and an inflammatory environment allows the establishment of these entities in the maternal system. The initial pro-inflammatory stage enables vasculature to develop, removes the cellular byproducts of implantation and results in the clinical manifestation of "morning sickness" that is seen in the early stages of pregnancy(Mor & Cardenas, 2010). In the second phase, the initial inflammatory state resolves and an anti-inflammatory state begins to predominate; creating a safe environment for fetal growth. In the final phase, a pro-inflammatory environment is induced in order to deliver the fetus (Romero et al., 2006). Increased inflammatory markers contribute to the promotion of the rupture of membranes, uterine contractions, and delivery (Mor & Cardenas, 2010).

### Cytokines

Cytokines are proteins produced by cells that are generally classified as proinflammatory or anti-inflammatory (Denney et al., 2011). There is redundancy of function between cytokines. Multiple cytokines illicit the same action, and have multiple target cells, so it can be difficult to attribute an action to a specific cytokine (Miyajima, Hara, & Kitamura, 1992). Cytokines are secreted by, and activate, various cells involved in the host immune response including activated phagocytes, epithelial cells, and T cells (Abbas, Lichtman, & Pillai, 2012). In general, cytokines produced by T-helper 1 (Th1) cells are pro-inflammatory and cytokines produced by T-helper 2 (Th2) cells are antiinflammatory and some cytokines exhibit properties of both (Brogin Moreli, Cirino Ruocco, Vernini, Rudge, & Calderon, 2012). An imbalance of pro versus antiinflammatory cytokines produces inflammation or muted immune responses. This can be attributed to the magnification of the normal synergistic or antagonistic effects that cytokines exhibit (Abbas, Lichtman, & Pillai, 2012). During normal pregnancy there is a shift to upregulate expression of *Th2* cells and suppression of *Th1* cells to prevent abortion of the fetus during development (Thellin & Heinen, 2003). Disruption of this altered balance of the immune system during pregnancy has been associated with gestational diabetes, preeclampsia, preterm labor, abortion and infection (Brogin Moreli et al., 2012; Fichorova et al., 2011). However, studies investigating levels of cytokines throughout pregnancy have presented dissimilar cytokine values during normal pregnancy (Curry et al., 2008; Makhseed et al., 2000; Vassiliadis, Ranella, Papadimitriou, Makrygiannakis, & Athanassakis, 1998). Variations in results could be explained by

different methodologies used to measure the cytokines and disparate sample sizes (30 versus 1200).

In a recent study, Fichorova et al. (2011) identified patterns in immune function markers that were specific to the type of bacteria present in the vaginal mucosa and placenta of pregnant women. They found that  $TNF-\alpha$ , IL-8 and ICAM-1 were elevated in the presence of *Gardnerella*, which is the most common causative pathogen of bacterial vaginosis. When multiple organisms associated with bacterial vaginosis were present, pro-inflammatory cytokine (TNF- $\alpha$ , IL-1 $\beta$ , IL-6), chemokine (IL-8), and acute phase marker (CRP and serum amyloid A) levels were elevated. Furthermore, *Lactobacillus*, which colonize the vaginal mucosa and are not pathogenic, suppress pathogenic strains and downregulate pro-inflammatory cytokines (Donato, Gareau, Wang, & Sherman, 2010; Othman, Neilson, & Alfirevic, 2007; Zeuthen et al., 2010). However, patterns of immune function serum markers were not analyzed related to GBS colonization or infection in any identified studies. For this study, TNF- $\alpha$ , IL-6, IL-10 and vitamin D (25[OH]D) status were selected specifically for analysis because of their identified association with infectious diseases and action during pregnancy and because serum levels for these specific cytokines were available to for secondary analysis for this study.

*TNF-a*: TNF- $\alpha$  is a serum immune function marker of interest because it is involved with coordination of the cytokine cascade and regulation of macrophage biology, which are both needed to fight infection. Alteration in TNF- $\alpha$  serum levels contribute to the development of various disease states, including sepsis and autoimmune conditions (Parameswaran & Patial, 2010). Macrophages are the primary producers of TNF- $\alpha$  in non-pregnant populations. However, in pregnant women the placenta

contributes to increased TNF- $\alpha$  levels throughout pregnancy (Brogin Moreli et al., 2012). The elevated TNF- $\alpha$  levels increase insulin resistance, which could contribute to the development of hyperglycemia or gestational diabetes during pregnancy (Kirwan et al., 2002). There have been no human studies analyzing TNF- $\alpha$  in pregnant women related to GBS infection or colonization. However, in vitro experiments show an increase in TNF- $\alpha$  production in cells exposed to GBS (Berner et al., 2002; Mikamo et al., 2004). Additionally, TNF- $\alpha$  and IL-6 levels in mice increase systemically when inoculated with GBS when IL-10 production is decreased resulting in 60% mortality (Puliti et al., 2000). Because TNF- $\alpha$  levels increase in laboratory and animal studies as a result from GBS exposure, serum TNF- $\alpha$  levels may be increased in pregnant women colonized with GBS. Furthermore, due to the inverse relationship TNF- $\alpha$  and IL-6 have with IL-10 in response to GBS exposure in animal models, IL-6 and IL-10 will also be evaluated in this study.

*IL-6:* IL-6 is a serum immune function marker that is a pro-inflammatory cytokine involved in the acute phase in the immune response. IL-6 is elevated in amniotic fluid of pregnant women with premature rupture of membranes due to intrauterine infections and also increases during active labor (Santhanam et al., 1991). Elevated serum IL-6 levels have also been identified in other inflammatory conditions experienced during pregnancy, like preeclampsia (A. Sharma, Satyam, & Sharma, 2007). It is unclear if a similar elevation in IL-6 occurs in women with GBS colonization. However, IL-6 production does increases in vitro and in animal studies with exposure to GBS (Berner et al., 2002; Mikamo et al., 2004; Puliti et al., 2002). The increase in IL-6 in response to GBS exposure has not been verified or validated in human studies. Therefore, in this

study we will evaluate the level of serum IL-6 to determine if a similar increase in IL-6 production occurs in pregnant women in response to GBS colonization.

*IL-10:* IL-10 is a serum immune function marker that inhibits the synthesis of proinflammatory cytokines, such as IL-6 and TNF- $\alpha$ , and stimulates the production of B cells and their differentiation into antibodies (Vieira et al., 1991). Additionally, IL-10 levels vary during pregnancy and may be involved in the maintenance of a viable pregnancy (Denney et al., 2011; Hashii et al., 1998). Early in pregnancy IL-10 is protective because it inhibits secretion of inflammatory IL-6, TNF $\alpha$ , and INF- $\gamma$  allowing the fetus and placenta to develop without being rejected by the maternal system. As the pregnancy progresses, the level of IL-10 decreases and the resulting increase in inflammatory cytokines allows the initiation of labor (Brogin Moreli et al., 2012).

Reduced IL-10 levels are associated with fetal loss in the first trimester, preeclamsia, gestational diabetes, and preterm birth (Brogin Moreli et al., 2012). Serum levels of IL-10, as related to GBS colonization in pregnant women, have not been evaluated. Although, Madureira et al. (2011) conducted a study using a murine model and found reduced levels of IL-10 in animals that carry GBS antibodies, which conferred immunity to the bacteria in offspring. Conversely, Bebien et al. (2012) found that the βh/c component of GBS induced IL-10 production. The higher production of IL-10 inhibited IL-12 production, which is involved in inducing immune responses, resulting in GBS being able to escape host cell detection and survive. When Bebien administered recombinant IL-10 to the GBS infected mice to test the effect of IL-10, the number of GBS in the mice increased significantly. If results from the animal studies described can be translated to humans, it is expected that there will be some elevation in serum IL-10

levels later in pregnancy in women colonized with GBS. Elevated IL-10 levels could be the result of GBS presence, like Madureira's study, or because the normal IL-10 elevation in early pregnancy enables GBS colonization in exposed women, like Bebien's study. Regardless of the mechanism, elevated IL-10 could be a clinical laboratory indicator for GBS colonization in pregnant women during the third trimester. *Vitamin D (25[OH]D) Status* 

# Circulating vitamin D (25[OH]D) deficiencies have been associated with susceptibility to infectious diseases such as influenza, tuberculosis, and pneumonia (Chesney, 2010). Vitamin D3 is the form of vitamin D that is produced by the skin in response to sunlight and primarily the form of vitamin D consumed from foods containing vitamin D. Vitamin D3 is then converted to Vitamin D (25[OH]D) in the liver. Vitamin D (25[OH]D) is the circulating form of the vitamin that is traditionally measured to identify vitamin D status in clinical populations (Hollis, 2005, 2008, 2012). It is generally accepted that serum vitamin D (25[OH]D) levels must be above 20 ng/ml to maintain normal physiologic processes and fetal development during pregnancy. Recently, vitamin D (25[OH]D) experts suggest that levels greater than 32 ng/ml are necessary to support all physiologic processes that require vitamin D (25[OH]D) for optimal functioning, like preventing infections (ACOG, 2011; Holick, 2011; Hollis, 2012). Therefore, the normal increase in serum vitamin D (25[OH]D) levels during pregnancy may be involved in preventing infection and colonization with pathogenic bacteria during normal pregnancy. There is a possibility that colonized women may have lower serum vitamin D (25[OH]D) levels, predisposing them to group B streptococcus colonization. To date, no randomized control trials evaluating the effects of vitamin D

(25[OH]D) supplementation during pregnancy related to maternal colonization or infection have been completed (De-Regil Luz, Palacios, Ansary, Kulier, & Peña-Rosas Juan, 2012). However, a meta- analysis conducted by Thorne-Lyman and Fawzi (2012) suggests that it is unknown how vitamin D (25[OH]D) relates to maternal infections since the relationship between vitamin D (25[OH]D) and immunity has only recently been established. There is currently no documented relationship between serum vitamin D (25[OH]D) levels and maternal GBS colonization.

# Implications for the Nursing Discipline

Person, health, environment, and nursing are the metaparadigm concepts that remain the pillars of the nursing discipline. Perception of these concepts constantly evolves to incorporate new knowledge gained through practice, research, education, and exposure to other disciplines. Since most human disease processes are multifactorial in nature and nurses interact with individuals throughout the illness-wellness continuum, it is imperative nurses understand how gene-environment interactions impact health. Research investigating exogenous and endogenous, like DNA methylation and serum cytokines and how they respond to environmental exposures, continues to generate data that improves our understanding of factors associated with complex disease processes. As the state of the science continues to evolve, nurses must begin to incorporate new data into their own research to deliver the best possible care to patients.

Investigations designed to identify biologic mechanisms explaining how epidemiologically defined risk factors result in complex disease processes lag behind data generated by epidemiologic studies (Dempfle et al., 2008; Hunter, 2005; Khoury, Davis, Gwinn, Lindegren, & Yoon, 2005; Martino & Prescott, 2011). Nurses, as members of

interdisciplinary teams, can utilize advances in epigenomic techniques to better assess levels of health and disease risk that may help explain findings identified in epidemiological studies. Nurses are well suited to investigate factors that may contribute to epigenomic variation because by nature of the profession, nurses continuously bridge science and technology to patient populations (Clark, Adamian, & Taylor, 2013; Loescher & Merkle, 2005). This study aimed to identify DNA methylation patterns and serum immune system markers associated with maternal GBS colonization. If vitamin D (25[OH]D) plays a role in the modulation of serum immune system markers, it would be a cost effective clinical intervention that could be introduced into practice to reduce GBS colonization rates and an alternative to antibiotic treatment. This study exemplifies how translational research can be initiated by hypotheses from the bedside, examined at the bench and brought back to the bedside to improve health outcomes.

### CHAPTER III

### **METHODS**

### Design

An exploratory secondary data analysis was completed using acquired quantitative data and maternal peripheral blood samples that were previously collected in a prospective longitudinal cohort study of nulliparous pregnant women. The data was initially collected to evaluate differences in women with and without preeclampsia. All women who had preeclampsia were excluded from this analysis because women with preeclampsia have different methylation patterns than women without preeclampsia (Anderson, Ralph, Wright, Linggi, & Ohm, 2013). The participants were enrolled in the primary study during the first trimester of their pregnancy and were followed through the time of childbirth. Eligibility criteria for the primary study included; English speaking, no previous births after 20 weeks gestation, age >18 years, and singleton pregnancy. At enrollment between 10-14 weeks gestation, baseline demographic information and venipuncture to collect blood for genome-wide DNA methylation analysis and other serum blood tests including TNF-a, IL-6, IL-10 and circulating vitamin D (25[OH]D were completed. In subsequent trimesters (second trimester 22-26 weeks gestation and third trimester 32-36 weeks gestation), venipuncture to collect blood for serum blood tests including TNF-α, IL-6, IL-10, and circulating vitamin D (25[OH]D) was completed. A subgroup of 6 women (n=2 GBS positive; n=4 GBS negative) were selected as control

samples for genome-wide DNA methylation analysis in the primary study. The methylation analysis was completed on peripheral blood samples obtained in the first trimester using the Infinium bead-based array platform (Illumina, Inc., San Diego, CA). For this study, only women (n=42) with documented uncomplicated (no preeclampsia, gestational diabetes, chorioaminoitis, etc) pregnancies in the primary study were included in the cytokine and vitamin D analysis. A subset of 18 women (n=9/group) were evaluated for differential genome wide methylation differences.

### Procedures

### Collection of Physiologic Data

Prior to commencement of data collection, this study was approved by the University of North Dakota Institutional Review Board (Appendix A). The primary study was approved by the University of North Dakota Institutional Review Board (Appendix B) and Altru Health System (Appendix C) prior to enrollment of participants into the study. Informed consent was completed by participants after verbal and written descriptions of the parent study were given. Participants were informed that blood samples would be taken and laboratory studies would be conducted on DNA extracted from these samples. Serum collected for cytokine and vitamin D analysis was collected via venipuncture. Whole blood was collected in a red top vacutainer (B-D) blood tube, retained at room temperature and allowed to clot for 1 hour followed by centrifugation at 1,000 x g at 4 degrees centigrade for 10minutes. Serum (500 µl) was placed in separate vials and frozen at -80 degrees centigrade until they were analyzed for the parent study. Cytokine laboratory analysis were completed at the United States Department of Agriculture, Agricultural Research Service (USDA ARS) Grand Forks Human Nutrition Center and vitamin D laboratory analysis was completed in the Brown-Borg laboratory at the University of North Dakota, School of Medicine and Health Sciences for the parent study. Only de-identified data was utilized in the current study. Medical history, GBS colonization status, and information about the participants had previously been extracted from the medical record for the purpose of the primary study using a standard data abstraction form (Appendix D). Descriptive statistics were used to elicit information about the study population including: age, race/ethnicity, ethnicity, weight, sex of the infant, gestational age at birth, and co-morbidities.

# Cytokine Laboratory Analysis

Cytokine analysis that was previously completed on serum samples for TNF- $\alpha$ , IL-6, and IL-10 (Bio-Plex, Millipore, Fountain Hills, AZ) at each of the three pregnancy trimesters. In brief, the target protein antibody was coupled to dual beads and incubated with sample. The protein of interest was captured, combined with a biotinylated antibody for a different epitope, and detected using a dual-laser flow based reader. For the cytokine analyses, the serum was diluted one volume of sample to three volumes of Bio-Plex human serum sample diluent. Next, 50  $\mu$ l of assay diluent was added to each well in the 96 well plate. Then, 200  $\mu$ l of cytokine standard, control, or sample was added to each plate and allowed to incubate at room temperature for two hours. All fluid from the wells was aspirated and each well washed a total of four times. 200  $\mu$ l of conjugate was then added to each well and allowed to incubate a room temperature for two hours. After aspirating all fluid from the wells and washing the wells four times, 200  $\mu$ l of substrate solution was added to all the wells, covered with foil and incubated at room temperature

for 20 minutes. Lastly, 50  $\mu$ l of stop solution was added to each well and read the plate at 450nm within 30 minutes with the wavelength correction set at 540 or 570nm.

### Vitamin D (25[OH]D) Laboratory Analysis

Circulating vitamin D (25[OH]D) levels were previously analyzed from serum as follows using the Immunodiagnostic Systems Ltd (IDS) 25-Hydroxyvitamin D enzymeimmunoassay (EIA) kit. Per the manufacturers assay protocol, the assay allows for 39 samples to be run in duplicate with 2 built in controls. The assay has good correlation with other methods for determining vitamin D (25[OH]D) in both serum or plasma with excellent sensitivity and specificity (5.3 - 7.4%) variability within assay, and 5.3 - 11.7% between assays (Hyppönen, Turner, Cumberland, Power, & Gibb, 2007). Briefly, the procedure for vitamin D (25[OH]D) EIA analysis was completed by adding 25 µl of serum to 1 ml of a propriety buffer reagent that dissociates vitamin D from protein. The diluted samples were then incubated at room temperature in a vitamin D antibody coated plate for 2 hours. Enzymes that bind selectively to the vitamin D biotin complex was then added and the samples were then washed. Next, a chromogenic substrate was added to the samples. The reaction was then stopped by adding a hydrochloric acid solution so the intensity can be measure of the treated vitamin D biotin labeled complex using a microtitre plate reader.

### DNA Methylation Laboratory Analysis

To complete the DNA methylation analysis portion of this study, the following steps were completed to examine the hypothesis that "differences in maternal DNA methylation patterns during early pregnancy are present in women with and without GBS colonization." DNA methylation was previously quantified in peripheral white blood

cells, collected in the first trimester, in a subset of six participants in the primary study using the Infinium bead-based array platform at the University of Minnesota's Biomedical Genomics Center and Illumina Core laboratory. Of those participants, four screened negative for GBS and two screened GBS positive in the third trimester.

In order to identify trends and potential differences in methylation between GBS positive and negative women, 12 additional samples were analyzed for this study (n=5 GBS negative; n=7 GBS positive). Samples were randomly selected from the remaining samples from the primary study. Genome-wide DNA methylation was determined using the Infinium platform (Illumina, Inc., San Diego, CA) at the University of Minnesota's Biomedical Genomics Center and Illumina Core Laboratory. Results from these samples were pooled with the results previously obtained to evaluate changes in DNA methylation between groups using the GenomeStudio Data Analysis Software that is compatible with output from Illumina. DNA methylation analysis was also completed using R 3.0.2 Statistical Environment for Windows.

DNA methylation at over 485,000 individual CpG dinucleotides in peripheral blood collected from women with and without GBS colonization (n=9 GBS negative; n= 9 GBS positive) was quantified. Samples underwent bisulfite conversion, which turns unmethylated cytosines to uracil and leaves methylated cytosines unaffected. The change allowed two query probes to detect differences in methylation between samples for predefined segments of DNA based on binding of specific nucleotides. This process is known to have greater than 99% conversation efficiency, yielding highly sensitive singlenucleotide resolution of methylation status. The quantitative amount of methylation at each CpG loci was reported as a beta value ( $\beta$  = methylated sites / (unmethylated gene +

methylated sites + 100)) (Bibikova et al., 2011). Beta values range from 0-1 and represents the percentage of DNA methylation (0-100) present at a given site. The Infinium platform processes 12 samples per plate.

Individual CpG dinucleotides that differed in DNA methylation between women with and without GBS colonization by +/- 20% as a percentage of total methylation, were designated as differentially methylated. Statistical significance of these changes between groups was then determined by *t*-tests (using a two-tailed, alpha of 0.05) (see Data Analysis section). In order to determine the potential for clinical relevance of differentially methylated CpG sites, functional analysis of genes with differentially methylated CpGs was also completed (see Functional Analysis section).

The specific function of individual genes was verified utilizing GeneCards and Pubmed. In order to validate the array, six genes were selected for validation of methylation patterns via bisulfite sequencing by Genwiz, Inc. (South Plainfield, NJ), based on greatest mean methylation differences. Two of the six gene sites also underwent a cloning step to assist in identifying methylation on specific CpG sites since the quality of DNA may not have been sufficient to obtain accurate reading without magnification because it is a clinical blood sample with multiple cell types present. After the samples were bisulfite treated, sequencing was used to determine the DNA sequence and validate the results from the Illumina Infinium array. The next sections describe the process in greater detail.

### Extraction of White Blood Cells from Blood

For the initial study, whole blood was obtained via venipuncture and collected in a purple top vacutainer (B-D) blood tube containing ethylenediaminetetraacetic acid

(EDTA). Collection tubes were immediately placed on ice during transport to the GFHNRC, followed by centrifugation at 1,000 x g at 4°C for 10 minutes. Plasma was removed and buffy coat collected and aliquoted into 1ml vials and frozen at -80 degrees centigrade until they were analyzed. The WBCs separate from the rest of the sample into a "buffy coat" layer as a result of being mixed with the EDTA. All other cell types are lysed from the EDTA solution and the WBCs form a pellet in the bottom of the tube when centrifuged. After centrifuging, the liquid components above the pellet were pipetted off leaving the WBC pellet. The remaining debris was washed from the WBC pellet by resuspending the pellet in EDTA solution, centrifuging, and discarding the fluid. *DNA Extraction from WBCs* 

Pelleted cells from the previous step were resuspended and homogenized by adding 270  $\mu$ l of lysis buffer to the sample in preparation for DNA extraction. Next, 30 $\mu$ l of proteinase K (10 mg/ml from Invitrogen) was added to each sample tube and the tubes were vortexed to thoroughly mix the samples. The samples were then incubated overnight at 60° C. The next day, the samples were incubated at 100°C to deactivate the enzyme. The samples were then transferred into 2ml phase lock microcentrifuge tubes and 300  $\mu$ l phenol:choloroform:isoamyl alcohol (25:24:1) was added to each sample. The tubes were then centrifuged and the upper aqueous phase was transferred into a fresh microcentrifuge tube. Next 600  $\mu$ l of 100% ethanol and 30  $\mu$ l of 7.5 M ammonium acetate were added to each tube containing the aqueous phase and samples were incubated for 3 hours at -20°C. The samples were then centrifuged, the fluid was discarded and the remaining pellet was washed with 300  $\mu$ l of 75% ethanol. The samples were centrifuged again and the remaining DNA pellets were allowed to air dry.

### DNA Quantification

All of the DNA sample pellets from the previous procedure were then individually re-hydrated in 100  $\mu$ l of nuclease free water and the DNA concentrations of each sample were determined using the Epoch micro-volume spectrophotometer system. After each well of the spectrophotometer was calibrated by running all 16 plate wells with 2  $\mu$ l DNAse and RNase free water, 2  $\mu$ l of a DNA sample was added to each well to determine the amount of DNA in the sample. For this study, 30 $\mu$ l of each of the 12 samples were plated and sent to the University of Minnesota for genome wide methylation analysis. The remaining samples volumes were used to validate DNA methylation patterns identified by the Illumina platform.

### Bisulfite Conversion of DNA

To verify the DNA methylation patterns identified by the Illumina Infinium array, remaining DNA from six participants (3 GBS positive and 3 GBS negative) were treated for bisulfite conversion using the EZ DNA Methylation Kit (Zymo Research). Per the kit specifications, the protocol has greater than 99% conversion efficiency converting unmethylated cytosine residues to uracil and greater than 99% protection of methylated cytosines. Samples were prepared, based on the concentration of DNA in each sample after being quantified using the Epoch micro-volume spectrophotometer system, for an input DNA amount of 500ng/sample. For example, if the amount of DNA in a sample was 59.9 ng,

500 ng  $(1\mu l/59.5ng) = 8 \mu l$  of the sample with DNA was used for next step

The appropriate volume of DNA from each sample for 500 mg/sample along with 5  $\mu$ l of M-dilution buffer and nuclease free water was added to a microcentrifuge tube to

yield a total volume of 50  $\mu$ l for each sample. For example, 8 $\mu$ l of DNA sample + 5  $\mu$ l M-dilution buffer + 37  $\mu$ l water = 50  $\mu$ l. Samples were then incubated for 15 minutes at 37° C. Next, 100  $\mu$ l of CT conversion reagent was mixed with each sample and the samples were then incubated in the dark for 14 hours at 50° C.

The samples were then incubated at 0-4°C (on ice) for 10 minutes. While the incubation was ongoing, 400 µl of M-binding buffer was added to a Zymo-spin IC column tube for each of the six samples, then the samples were added to the Zymo-spin IC column tubes and mixed by inversion after incubation was complete. The tubes were then centrifuged at full speed ( $\geq 10,000 \text{ xg}$ ) for 30 seconds. The flow-through was discarded and 100 µl of M-wash buffer was added and the tubes were again centrifuged at full speed for 30 seconds. Next, 200 µl of M-desulphonation buffer was added to each sample tube and the samples were incubated at room temperature for 15-20 minutes. The samples were then centrifuged at full speed for 30 seconds and then washed with 200 µl of M-wash buffer. The samples were then centrifuged at full speed for 30 seconds and washed again with 200  $\mu$ l of M-wash buffer and centrifuged at full speed for 30 seconds. The Zymo-spin IC columns for each sample were then placed into a fresh 1.5 ml microcentrifuge tube and 10 µl of M-elution buffer was added directly to the column matrix and centrifuged for 30 seconds. The samples were then stored at -20°C until the primers were designed to complete polymerase chain reaction (PCR) and validation sequencing.

#### Primer Design

Primers were designed to cut the DNA and validate methylation patterns in six sites identified as differentially methylated between women with and without GBS colonization. The sites were selected based on greatest difference in methylation between groups that were known genes (3 sites with loss of methylation, 3 with methylation gain). Two of the six sites were also be amplified using TOPO cloning reactions in anticipation that the amplification of the DNA samples via PCR alone may not yield adequate quality samples for sequencing. The DNA sequences for the six sites were determined utilizing University of California, Santa Cruz (UCSC) Genome Browser (Kent et al., 2002). The chromosome number and map information for the genes, from the Illumina Infinium output, were entered into the search box. After selecting "DNA" from the "view" dropdown a new window opens allowing selection of how many nucleotides away from the CpG of interest you would like the software to search for a suitable region to cut the DNA for sequencing. Segments 250 bases upstream and 250 bases downstream were entered as criteria to ensure the region of interest would be in the resultant product after PCR was complete. The actual DNA sequence was then copied and pasted into The Li Lab Department of Urology, University of California, San Francisco MethPrimer design tool (Li & Dahiya, 2002). The identified primer sequence was then used to order the primers from Integrated DNA Technologies (IDT) (http://www.IDTDNA.com) and utilized in the PCR step. DNA site sequences, primer sequences and properties of the primers supplied by IDT are located in Appendix E. When the primers arrived from IDT, all were rehydrated into a stock solution of  $1\mu g/\mu l$ . For example, one vial was 0.27 mg/vial, therefore 270 µl of DNAse/RNAse free water was added to rehydrate the

sample. From the stock solutions, a working concentration of 200 ng/ $\mu$ l for each primer was created by adding 20  $\mu$ l of stock and 80  $\mu$ l of DNAse/RNAse free water to a 1.5 ml microcentrifuge tube. Stock and working concentrations for each primer were then stored at -20°C.

#### Polymerase Chain Reaction

To complete the PCR step for each primer, the  $ZymoTaq^{TM}$  PreMix protocol was completed using the bisulfite treated DNA (described previously) from each of the six participants (3 GBS positive, 3 GBS negative) for all six primers. The following reaction set up was used for each primer for each sample of DNA: 25 µl *Zymo Taq^{TM*} PreMix, 1 µl forward primer, 1 µl reverse primer, 1 µl bisulfite treated DNA, 22 µl DNase/RNase free water. The following conditions were used for the PCR reaction using a hot start: initial denaturation at 95°C for 10 minutes, followed by 35 cycles of denaturing at 95°C for 30 seconds, annealing at 54°C for 35 seconds, and extension at 72°C for 45 seconds. After the 35 cycles were completed, final extension at 72°C for seven minutes was done followed by a hold at 4°C for 30 minutes. The PCR products were visualized by electrophoresis in agarose gel (0.5 gm agarose, 50ml TAEx1 buffer, with 5µl DNA star) at 100V with a 100bp ladder. 13.5 µl of each sample was loaded into the gel with 1.5 µl x10 loading dye. Products were produced for all primers (Appendix F).

#### Methylation Validation

In order to validate the methylation patterns identify by the Illumina Infinium array, the PCR products that were produced in the previous step were sent to Genewiz, Inc. (South Plainfield, NJ) for sequencing. A portion of the PCR products from ANXA2

and RHPN1 were cloned using the Topo TA-cloning kit; all remaining products were purified using the QIAquick® PCR purification kit described in the following sections.

*PCR Clean-up:* First, five volumes of PB buffer were added for every volume of PCR product for each sample. The samples were then transferred to QIAquick spin column tubes and centrifuged for 60 second. After discarding the flow through, the samples were washed with 0.75ml of PE buffer, then centrifuged 60 seconds. After removing the flow through, samples were centrifuged again for 60 seconds and the columns were then placed in a clean 1.5 ml microcentrifuge tube. Then, 50  $\mu$ l of EB buffer was added to the center of the column and centrifuged for one minute to elute the DNA. All samples were then sent to Genewiz, Inc. (South Plainfield, NJ) for sequencing (Appendix G). Since the samples were not of adequate quality for sequencing, cloned samples were also sent to validate the array.

*TOPO TA-cloning:* PCR products from ANXA2 and RHPN1 were used in the cloning protocol to optimize segments for sequencing using the TOPO TA Cloning Kit for Sequencing (Invitrogen, Life Technologies). Briefly, 4  $\mu$ l of fresh PCR product was incubated at room temperature for 30 minutes with 1  $\mu$ l salt solution (1.2M NaCl, 0.06 M MgCl<sub>2</sub>), and 1  $\mu$ l TOPO vector, then placed on ice. Next, 2  $\mu$ l from the completed reactions were added to a vial of One Shot® Chemically Competent *E. coli*, which were then incubated on ice for 30 minutes. The samples were then heat shocked at 42 degrees C for 30 seconds, then placed back on ice. Next, 25  $\mu$ l of room temperature Super Optimal broth with Catabolite repression (S.O.C. medium) was added to each sample and the samples were shaken horizontally at 200 rpm for one hour at 37 degrees C. Kanamycin selective agar plates were prepared and two different concentrations (20  $\mu$ l

and 50  $\mu$ l) were plated for each sample to ensure adequately spaced colonies. The plates were then incubated at 37 degrees C overnight. Then, 10 white colonies were selected and suspended in individual test tubes with 3ml Luria Broth (LB broth) containing 50  $\mu$ g/mL kanamycin and incubated overnight.

The plasmid DNA was then extracted using the QIAprep® Spin Miniprep Kit High-Yield Protocol. First, sample tubes were centrifuged at 8,000 rpm for three minutes at room temperature to pellet the bacteria with plasmid products. The pellets were then resuspended in 250 µl P1 buffer and placed into a microcentrifuge tube. Next P2 buffer was added to the samples and mixed by inverting the tube 10 times. Then 350 µl of N3 buffer was added to each tube, mixed by inverting 10 times and then centrifuged at full speed for 10 minutes. The supernatant was then pipetted into QIAprep spin columns and centrifuged at full speed for 60 seconds and the flow through was discarded. Next, the samples were washed with 500 µl PB buffer and centrifuged again at full speed for 60 seconds. Then 750 µl of PE buffer was added to each sample and centrifuged for 60 seconds. Samples were centrifuged for an additional minute to ensure all wash buffers had been removed. The spin columns were then placed into a clean microcentrifuge tube and 60  $\mu$ l of EB buffer was added to each tube. Samples were incubated at room temperature for 60 seconds and were then centrifuged for one minute to elute the DNA. Samples were then sent to Genewiz, Inc. for validation; and sequencing was successfully matched (Appendix G).

#### Data Analyses

#### Statistical Analysis

All cytokine and vitamin D (25[OH]D) data analysis procedures were performed using Statistical Package for Social Sciences (SPSS) version 22. All variables were examined for violations of statistical assumptions, including missing values and outliers, with SPSS Frequencies, and Explore. Upon completion of data screening to ensure all univariate and multivariate assumptions were met, descriptive statistics were completed to describe the sample characteristics and compare the groups (GBS positive and negative). The normality of the distribution of cytokines was determined by evaluating the skewness, kurtosis, and Kolmogorov-Smirnov statistics for each variable. Pearson correlation tests were completed to determine if cytokines and vitamin D (25[OH]D) levels co-varied throughout pregnancy. Since there was no significant correlation among these variables, a series of repeated measures ANOVAs were performed to compare each of the three cytokines (TNF- $\alpha$ , IL-6 and IL-10) and vitamin D (25[OH]D) between the two groups (n=16 GBS positive and n=26 GBS negative) from each of the three trimesters of pregnancy. Mauchly's test was used to identify violations in sphericity and bonferroni corrections were applied for multiple comparisons. The Greenhouse-Geisser correction for a violation in sphericity was used for vitamin D (25[OH]D) analysis. Given the restrictions of performing a secondary analysis, it was not feasible to change the sample size. However, performing a compromised post-hoc power analysis using "G\*power" software indicated that the current sample size allowed 69% power and a medium effect size of 0.3. This particular type of power analysis was developed for studies where the sample size cannot be altered and equates the risk of committing a

Type I and Type II error (Faul, Erdfelder, Buchner, & Lang, 2009; Faul, Erdfelder, Lang, & Buchner, 2007). This approach to analysis helps in identifying trends and effect size so sample size can be determined in future investigations that have adequate power.

Mean values for serum cytokine and vitamin D (25[OH]D) levels were utilized to conduct a one-way multivariate analysis of variance (MANOVA) to correct for confounding resulting from data being collected over time, determine observed effect size and observed power. The results obtained were then used in G\*power to determine the number of participants that would be required to see an effect with the power and effect size calculated from the study sample. All data analyses procedures in the study were performed using a two-tailed alpha of 0.05.

For DNA methylation analysis, fluorescent background intensities were normalized using the Genome Studio software. A series of negative controls are embedded into the assay that the software utilized to generate the detection p values for each probe. The specific normalization process applied when using GenomeStudio software is propriety information, and a limitation of using this software for methylation analysis (Gentleman et al., 2004; Hansen et al., 2013, Smyth, 2005). Probes that had detection *p* values greater than 0.05 were considered not to have reliable signal intensities and eliminated from analysis. Independent sample *t*-test comparisons were conducted to examine differences in mean DNA methylation at individual CpG sites between GBS positive and negative groups. The selection of independent sample *t*-test was driven by the fact the methylation testing was limited to 18 participants and DNA methylation was reported as a percentage of methylation at each CpG site. Findings from this analysis will inform the researcher of the effect size so that proper sampling can be determined for

future research. All statistical techniques were performed using a 2-tailed alpha of < 0.05 or 95% confidence interval. Statistical analysis of raw data provided by Illumina was also completed using the minfi package (Hansen, Ayree, & Irizarry, 2013) in the R programming environment (R Core Team, 2013) to first normalize the data and identify differentially methylated regions by using the limma package (Smyth, 2005). Significance testing was also completed using Benjamini and Hochberg's false discovery rate to correct for multiple comparisons (Benjamini & Hochberg, 1995) because it is the recommended standard and used for a majority of studies investigating specific clinical outcomes in human samples (Allison, Cui, Page, & Sabripour, 2006; Benjamini & Hochberg, 1995; Michels et al., 2013; Storey, 2003; Tusher, Tibshirani, & Chu, 2001; Wilhelm-Benartzi et al., 2013; York, 2003). R code used for analysis with normalization figures are located in Appendix I. Established work flows with detailed explanations of computations and code are available in open source, published workflows and user manuals for minfi (Hansen, Ayree & Irizarry, 2013) and limma (Smyth, 2005) packages.

Briefly, raw data were preprocessed using a series of minfi commands (Appendix I) to normalize the intensity for methylated and unmethylated channels (G. K. Smyth, Yang, & Speed, 2003). For the statistical analysis of DNA methylation, M-values were used because they are logit transformed  $\beta$  values and result in data that more closely follows a normal distribution and preferred for statistical methods such as a *t*-test. Beta values exhibit severe heteroscedasticity for extremes of methylation (highly methylated or unmethylated). Because the logit transformation corrects for this, M-values outperform beta values in terms of detection rate and true positives for detecting differences of CpG sites on the extremes of percent methylated. Dedeurwaerder et al.'s (2011) conversion

method for beta to M-values was utilized for analysis because a peak correction can be performed on the M-values to adjust the Infinium II probes. Infinium claims there are no differences between type I and II detection probes that affect detection of differential methylation (Bibikova et al., 2011). However, design I type probes have been shown to be more stable and have a more extensive detection range than type II probe, therefore require correction for statistical analysis (Dedeurwaerder et al., 2011). Using M-values results in more accurate analysis of the data, although are difficult to clinically interpret. Therefore, after using the logit transformed values (M-values) for statistical analysis, the data were back transformed to yield peak-corrected beta values for reporting and to allow for easier clinical interpretation of findings because beta values are biologically meaningful (Du et al., 2010). Sites with known single nucleotide polymorphisms (SNPs) were excluded from the analysis as these sites can disturb the accuracy of the Infiniuim probes (Bibikova et al., 2011b; Dedeurwaerder et al., 2011). Additionally, SNP sites were excluded to remove confounding effects that could result from having two different modifications at the same site. Furthermore, SNP analysis is beyond the scope of this project and the original consent form was not written to include this type of analysis.

After all the raw files were normalized to ensure the peaks of the Infinium II design bead type probes were comparable to the peak locations of the Infinium I bead bead type probes and converted to M-values, differential methylation analysis was conducted using the limma package. This package was selected because it performs additional normalization and offers commands that are ideal for use with small data sets. Essentially, Symth (2005) designed the program to strengthen analysis for small data sets by "borrowing information across genes" (p. 4) using empirical Bayesian methods to help

control for variance; which results in less false discoveries for small data sets than other programs. All normalization and analytics commands were designed for two color channel arrays, including the Infinium platform used for this study (Gentleman et al., 2004; Smyth, Yang, & Speed, 2003; Smyth, 2004; Smyth, 2005). The series of R commands fit each CpG site into a linear model using least squares fitting, then a contrast matrix was created to compare values between GBS positive and GBS negative women and in the final normalization command, the empirical Bayesian command incorporates the use of array weights to improve identification of CpG sites that are most likely to be different (Ritchie et al., 2006; G. Smyth, 2005). This correction for multiple comparisons increases the likelihood of identifying clinically meaningful results and reduces the chance of identifying false positives.

#### Functional Analysis

In order to evaluate the last research question, that differential DNA methylation in genes associated with immune function and inflammation, a functional analysis of genes associated with differential DNA methylation was completed using Database for Annotation, Visualization, and Integrated Discovery (DAVID) Bioinformatics Resource version v6.7 (Huang, Sherman, & Lempicki, 2009a; Huang, Sherman, & Lempicki, 2009b). Functional analysis reveals potential mechanistic underpinnings of GBS colonization or other clinically relevant information. Gene lists with significant differentially methylated CpG sites were uploaded into the DAVID v6.7 database. Sites with a gain of methylation in women with GBS colonization were uploaded as one group and those with a loss of methylation were entered as a separate group, to investigate changes in function based on differences in methylation at those sites. DAVID

determines functional categories of genes and classifies them with "high" stringency selected by the investigator for analysis (Huang, Sherman, & Lempicki, 2009a; Jiao et al., 2012). Using a classification stringency of "high" to determine functional annotation clusters identifies a "tight, clean, and smaller numbers of clusters" (Huang et al., 2009a, p. 47) that are more likely to be associated with biologically meaningful results. Output generated by the DAVID software also includes enrichment scores for each cluster, with higher numbers indicating there may greater involvement in the disease state being studied. Pathways with enrichment scores greater than 1.3 are likely most important in the functional or disease process, although lower scoring clusters could also offer insight into biological mechanisms associated with the disease process. Statistical calculations presented related to the function of the genes are determined utilizing conservative correction methodologies for multiple comparisons (Bonferroni; Benjamini, and Hochberg) (Huang et al., 2009). The specific function of individual genes was verified utilizing GeneCards® and Pubmed and reviewed in the discussion chapter.

#### CHAPTER IV

#### RESULTS

This chapter presents the results by first providing general descriptive statistics of the sample characteristics, followed by presentation of the statistical analyses that were completed to answer the research questions as presented in chapter one.

#### Description of Sample

A total of 42 women had an outcome of uncomplicated pregnancy in the parent study and were included in the analysis for this study. Among those women, 38% were GBS positive (n = 16) and 62% were GBS negative (n = 26), 9 participants from each group had DNA methylation analyzed. The sample population was primarily comprised of Caucasian women (84%), reflecting the demographics of the area from which the subjects were recruited. Race was self-identified by participants upon entry to the parent study. Other demographic data of the participants are presented in Table 6, which shows that no statistically significant differences were found (age, weight, gestational age at delivery, or infant gender) between GBS positive and negative women. Notably, there are no significant differences in co-morbidities, infections, or antimicrobial usage. Infection data and antimicrobial data were extracted from the medical record for usage at any point during the pregnancy because it may alter the vaginal microbial composition. Currently, there is no information in the literature describing how long and when vaginal microbiota returns to a woman's baseline composition or how it may contribute to GBS status.

Variable	Total Sample (N=42)	GBS + (n=16)	GBS – (n=26)	t /χ²	р
Maternal Age (years) [M(SD)]	26.6 (4.10)	25.1 (3.05)	27.5 (4.46)	-1.806	0.194
Maternal Prenatal Weight (pounds) [M(SD)]	168.7 (35.5)	161.2 (31.6)	173.3 (37.5)	-1.074	0.573
Gestational Age at Birth (weeks) [M(SD)]	39.1 (1.25)	38.7 (1.45)	39.2 (1.08)	-1.488	0.763
Race [n(%)] Caucasian Multi-racial Native Hawaiian/ Pacific Islander	38 (90.5) 3 (7.1) 1 (2.4)	14 (87.5) 1 (6.25) 1 (6.25)	24 (92.3) 2 (7.7) 0 (0)	1.679	0.432
Infant Gender [n(%)] Female Male Missing	22 (52.4) 19 (45.2) 1 (2.4)	12 (75) 4 (25) 0 (0)	10 (38.5) 15 (57.7) 1 (3.8)	5.480	0.065
Maternal Asthma [n(%)]	7 (16.7)	2 (12.5)	5 (19.2)	0.323	0.570
Infection [n(%)] None Respiratory Urinary Tract Chlamydia Herpes Tuberculosis Unknown	28 (66.7) 3 (7.1) 3 (7.1) 1 (2.4) 1 (2.4) 1 (2.4) 5 (11.9)	10 (62.5) 1 (6.2) 1 (6.2) 0 1 (6.2) 0 3 (18.8)	18 (69.2) 2 (7.7) 2 (7.7) 1 (3.8) 0 1 (3.8) 2 (7.7)	3.998	0.677
Antimicrobial [n(%)] Yes No	13 (31) 29 (69)	5 (31.2) 11 (68.8)	8 (30.8) 18 (69.2)	0.001	0.974

Table 6. Maternal Characteristics Compared between GBS Positives and Negatives

P indicates significance level based on a two-tailed alpha of 0.05

#### Serum Cytokine and Vitamin D (25[OH]D) Analysis

Research Question 1: Are serum levels of TNF- $\alpha$ , IL-6, IL-10, and vitamin D (25[OH]D) different in pregnant women with GBS colonization than pregnant women without GBS colonization?

Cytokines and vitamin D (25[OH]D) levels were not normally distributed for the 42 participants (Table 7). However, ANOVA is considered robust to departures in normality. Kepple and Wickens (2004) argue when sample size is greater than approximately 12 ANOVA analyses can be successfully completed if normality has been violated. Further, the laboratory values analyzed in this study were highly variable and were not normally distributed in similar studies that evaluated cytokine levels (Curry et al., 2008; Makhseed et al., 2000). Because cytokine levels are variable between individuals (not normally distributed, making it impossible to identify outliers) all participant data were included in the analysis in order to assess if evaluation of serum cytokine levels and vitamin D (25[OH]D) would be a useful clinically relevant measure

Table 7. Tests of Normality for Serum Cytokines and Vitamin D (25[OH]D)

Variable*	Skewness	Kurtosis	Kolmogorov- Smirnov			Sha	piro-	Wilk
			Stat	df	Р	Stat	df	Р
TNF-α	1.484	3.989	0.141	42	< 0.001	0.851	42	< 0.001
IL-6	4.999	26.755	0.478	42	0.036	0.898	42	0.001
IL-10	0.462	-0.059	0.107	42	< 0.001	0.273	42	< 0.001
(25[OH]D)	1.918	5.797	0.196	42	0.200	0.947	42	0.052

Stat = statistic

P = significance 2-tailed 0.05

\*mean laboratory values across all trimesters

 $df = degrees \ of freedom$ 

for predicting GBS colonization susceptibility. Serum cytokine and vitamin D (25[OH]D) levels were described by time during pregnancy with the following notations: T1 = first trimester (10- 14 weeks); T2 = second trimester (22-26 weeks); and T3 = third trimester (32-36 weeks). Correlation tests indicate vitamin D (25[OH]D) levels were not correlated with serum TNF- $\alpha$ , IL-6, or IL-10 levels during pregnancy, except T1vitamin D (25[OH]D) and T3 IL-10 (Table 8). Therefore, vitamin D (25[OH]D) was not used as a covariate and repeated measures ANOVA was completed on serum vitamin D (25[OH]D), TNF- $\alpha$ , IL-6 and IL-10 independently.

Variable	T1 Vitamin D		T2 Vita	min D	T3 Vita	min D
variable	(25[0		(25[0		(25[0	
-		<b>」</b> /	ί Ε			<b>」</b> /
	r	P	R	Р	r	P
T1 TNF-α	-0.111	0.483	-0.265	0.090	-0.277	0.076
T2 TNF-α	-0.001	0.994	-0.005	0.975	-0.101	0.523
T3 TNF-α	-0.018	0.091	-0.031	0.846	-0.067	0.674
T1 IL-6	-0.192	0.224	-0.140	0.375	-0.162	0.305
T2 IL-6	-0.224	0.154	-0.147	0.352	-0.186	0.238
T3 IL-6	-0.215	0.172	-0.162	0.306	-0.183	0.245
T1 IL-10	0.142	0.368	0.074	0.642	0.063	0.693
T2 IL-10	0.007	0.963	0.093	0.557	0.018	0.911
T3 IL-10	0.310*	0.046*	0.214	0.173	0.129	0.415

Table 8. Serum Vitamin D (25[OH]D) and Cytokine Correlations

*r* = *Pearson correlation* 

*P=significance 2-tailed 0.05* 

#### $TNF-\alpha$

TNF- $\alpha$  levels were compared between GBS positive (n=16) and negative (n=26) women across three trimesters of pregnancy. Mean TNF- $\alpha$  levels were calculated for each group in each trimester (Table 9, Figure 4a). A repeated measures ANOVA revealed no significant difference in TNF- $\alpha$  between GBS positive and negative women, *F*(2, 80) = 2.187, *p* = 0.119,  $\eta^2$  = 0.052 with an observed power of 0.453.

IL-6 levels were compared between GBS positive (n=16) and negative (n=26) women across three trimesters of pregnancy. Mean IL-6 levels were calculated for each group in each trimester (Table 10, Figure 4b). A repeated measures ANOVA revealed no significant difference in IL-6 between GBS positive and negative women, F(2, 80) = 2.991, p = 0.056,  $\eta^2 = 0.070$  with an observed power of 0.566.

Table 9. Serum TNF-α Levels

Table 10. Serum IL-6 Levels

	Mean	(SD)		Mean	(SD)
	GBS +	GBS -		GBS +	GBS -
T1 TNF-α	6.20 (2.77)	6.44 (1.92)	T1 IL-6	1.06 (2.98)	1.28 (5.71)
T2 TNF-α	6.21 (2.74)	7.41 (4.75)	T2 IL-6	1.67 (4.21)	1.04 (5.31)
T3 TNF-α	8.71 (6.19)	7.46 (2.54)	T3 IL-6	1.25 (3.58)	1.31 (6.70)

#### IL-10

IL-10 levels were compared between GBS positive (n=16) and negative (n=26) women across three trimesters of pregnancy. Mean IL-10 levels were calculated for each group in each trimester (Table 11, Figure 4c). A repeated measures ANOVA revealed no significant difference in IL-10 between GBS positive and negative women, F(2, 80) = 0.445, p = 0.642,  $\eta^2 = 0.011$  with an observed power of 0.120.

#### Vitamin D (25[OH]D)

Vitamin D (25[OH]D) levels were compared between GBS positive (n=16) and negative (n=26) women across three trimesters of pregnancy. Mean vitamin D(25[OH]D) levels were calculated for each group in each trimester (Table 12, Figure 4d). Mauchly's test indicated a violation in the assumption of sphericity, W(2) = 0.551, p < 0.05. Based

on a Greenhouse-Geisser correction for the sphericity violation, a repeated measures ANOVA revealed no significant difference in vitamin D (25[OH]D) between GBS positive and negative women, F(1.380, 55.218) = 0.882, p = 0.384,  $\eta^2 = 0.022$  with an observed power of 0.169.

Table 1	11. Serum	IL-10	Levels
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	Mean (SD)			Mean (SD)	
	GBS +	GBS -		GBS +	GBS -
T1 IL-10	4.23 (3.27)	4.23 (3.06)	T1 Vit. D	27.22 (9.36)	23.76(6.21)
T2 IL-10	5.29 (3.48)	4.35 (4.55)	T2 Vit. D	30.18(10.94)	25.14(6.42)
T3 IL-10	4.64 (4.56)	4.08 (2.85)	T3 Vit. D	30.08(14.09)	24.89(7.06)

a) TNF-α

b) IL-6

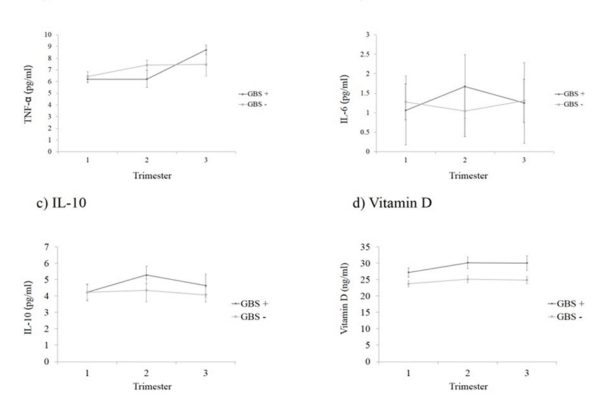


Figure 4. Cytokines and Vitamin D (25[OH]D) Across Pregnancy.

#### Multivariate Analysis

Mean values for serum cytokine and vitamin D levels were utilized to conduct a one-way multivariate analysis of variance (MANOVA) to correct for the confounding resulting from data being collected over time, determine effect size and observed power (Table 13). MANOVA revealed that serum cytokines and vitamin D levels do not significantly combine to affect GBS colonization status Wilks'  $\Lambda = 0.927$ , F(4, 37) = 0.725, p = 0.581, partial  $\eta^2 = 0.073$ . Further analysis comparing the sample based power and power needed to see a significant effect of cytokines and vitamin D on GBS status, indicates large sample sizes are required to see significant difference (Table 14)

	Mean ±	SD	Partial $\eta^2$	Observed Power
	GBS +	GBS -		
TNF-α (pg/ml)	$7.04 \pm 3.34$	$7.11 \pm 2.75$	< 0.001	0.051
IL-6 (pg/ml)	$1.33 \pm 3.57$	$1.21 \pm 5.89$	< 0.001	0.051
IL-10 (pg/ml)	$4.72\pm3.19$	$4.22 \pm 3.16$	0.006	0.077
(25[OH]D)(ng/ml)	29.16 ± 11.13	$24.59\pm6.24$	0.068	0.384

Table 13.	Multiva	riate Results
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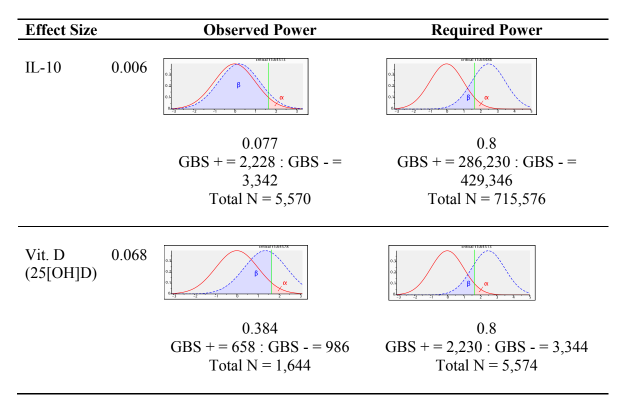


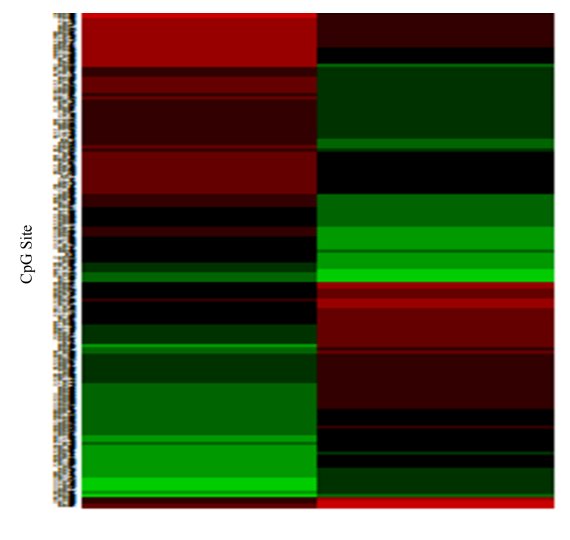
Table 14. Comparison of Sample Based Power to 80% Power with Required N

#### DNA Methylation Analysis

Research Question 2: Are DNA methylation patterns different between pregnant women with GBS colonization and those without GBS colonization?

#### GenomeStudio Statistical Analysis

Analyses comparing the pooled mean methylation of GBS positive and negative women (n = 9/group) identified a total of 141 CpG dinucleotides that were differentially methylated in maternal peripheral white blood cells between women who had a positive screening for GBS and those who had a negative screening. Figure 5, depicts the average beta scores of the 141 CpG dinucleotides that were identified as differentially methylated by greater than 20% betweem GBS positive and GBS negative women and statistically significant with a p value of less than 0.05. Of the 141 CpG sites, 62 sites had a gain of methylation and 79 sites had a loss in methylation in women with positive GBS screening compared to GBS negative women. CpG sites with differential methylation were distributed across the chromosomes (Figures 6a and 7a), associated with known genes (62%) and located predominantly in the body (66%) of genes (Figures 6b and 7b). Approximately half of the CpG sites (49%) were not associated with CpG islands (Figures 6c and 7c).

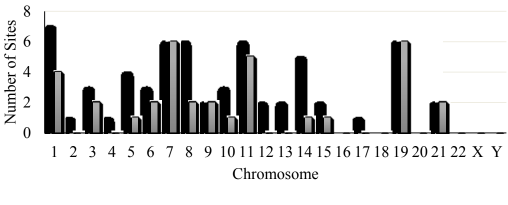


GBS Negative

**GBS** Positive

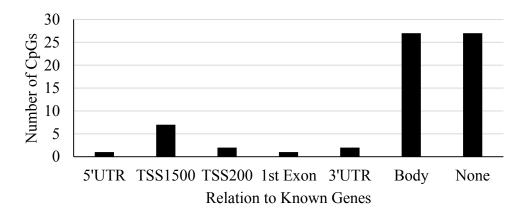
Figure 5. Differential Methylation with GBS Colonization (GenomeStudio).

a) Chromosomal Distribution



■ Total CpGs ■ Known Genes

b) Proximity to Gene



c) Proximity to CpG Island

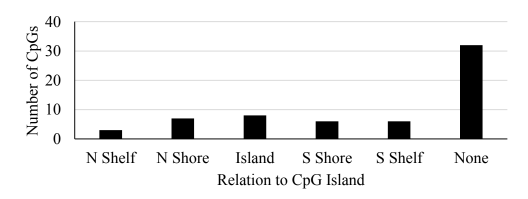
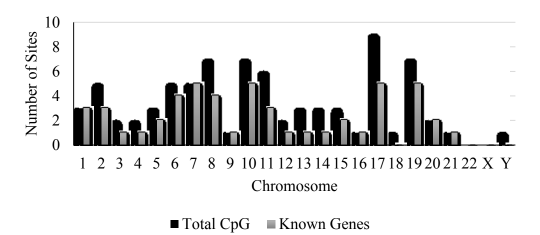
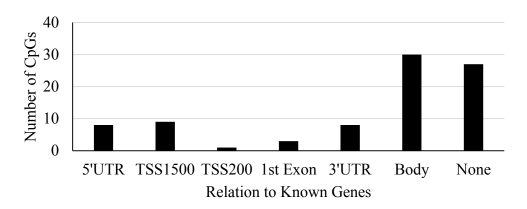


Figure 6. DNA Methylation Gain Distributions with GBS (GenomeStudio).

a) Chromosome Distribution



b) Proximity to Genes



c) Proximity to CpG Island

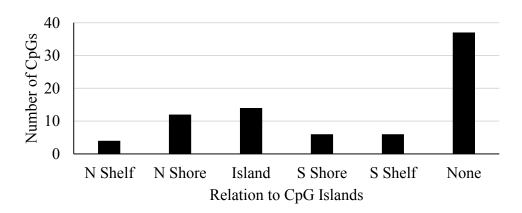


Figure 7. DNA Methylation Loss Distributions with GBS Colonization (GenomeStudio).

#### *R* Statistical Environment Analysis

Analyses conducted in R comparing the pooled mean methylation of GBS positive and negative women (n = 9/group) identified no statistically significant differences in methylation at any CpG site when a 20% change in beta value when FDR was used to determine significance at any level is used for analysis (adjusted p=0.001, 0.05, 0.01). Figure 8 is a volcano plot presenting all CpG sites with parameters set to highlight blue any sites with greater than a 20% difference in methylation between GBS positive and GBS negative women that met significance using an FDR of 0.05. No blue CpG sites are present on the graph, indicating there is no statistically significantly different methylation at any CpG site by GBS status for this sample (R code and mathematical code returning zero results available in Appendix I). The plot is designed using the log values for more concentrated clear visualization of the data. In order to compare the results using the R programming environment to the results produced using the GenomeStudio software, the analysis was also conducted using the same unadjusted p value to determine significance at 0.05. A total of 125 CpG dinucleotides that were identified as having greater than 20% difference in methylation in maternal peripheral white blood cells between GBS positive and negative women. Of the 125 CpG sites, 54 sites had a gain in methylation and 71 sites had a loss in methylation for women with positive GBS screening at 37 weeks compared to GBS negative women when alpha is 0.05. CpG sites with differential methylation were distributed across the chromosomes (Figures 9a and 10a), associated with known genes (62%) and located predominantly in the body (59%) of genes (Figures 9b and 10b). Approximately half of the CpG sites (51%) were not associated with CpG islands (Figures 9c and 10c).

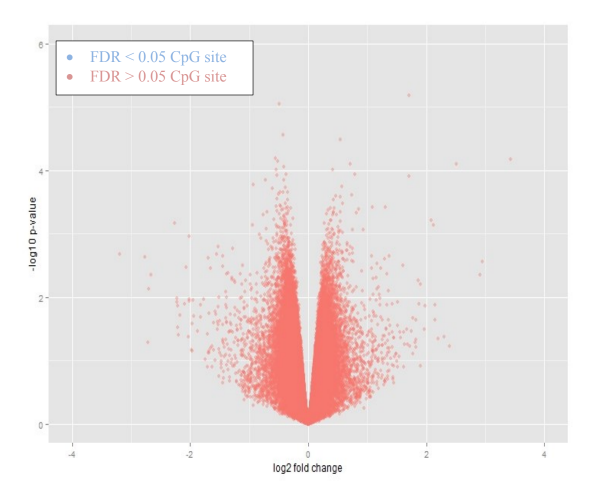
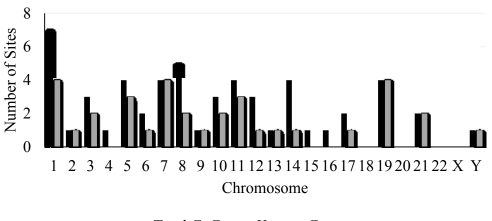


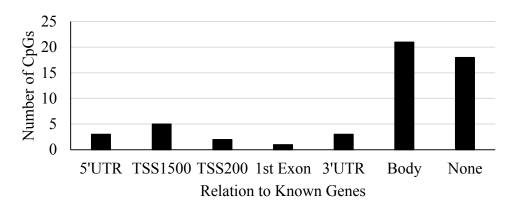
Figure 8. Volcano Plot of CpG Methylation Differences by GBS Status (R)

a) Chromosomal Distribution



■ Total CpGs ■ Known Genes

b) Proximity to Gene



c) Proximity to CpG Island

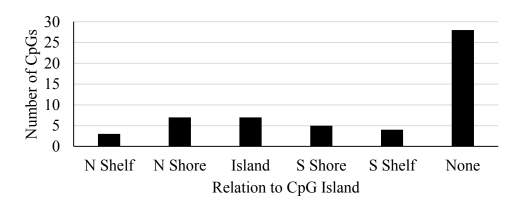
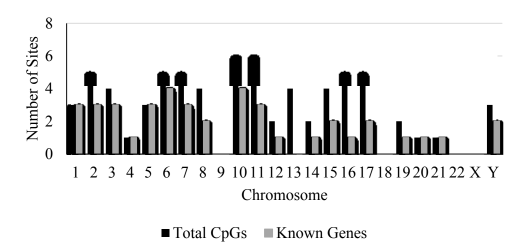
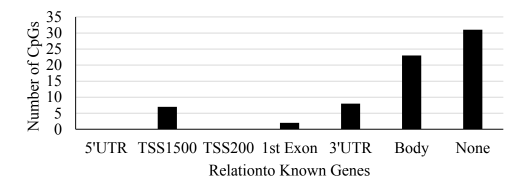


Figure 9. DNA Methylation Gain Distributions with GBS (R).

a) Chromosomal Distribution



a) Proximity to Gene



b) Proximity to CpG Island

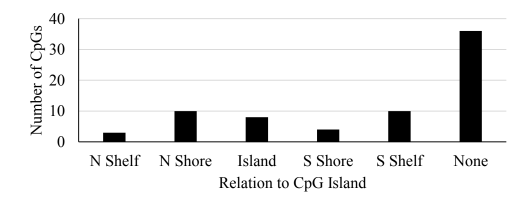


Figure 10. DNA Methylation Loss with GBS (R).

### Gene Comparison

Using two different statistical software applications to identify significant differentially methylated CpG sites yielded a discordant number of CpG sites, 141 using GenomeStudio software and 125 using R programming environment. Of the known genes associated with the CpG sites, 52% of the genes identified were common in the results using both packages, 29% were specific to analysis with the GenomeStudio platform and 19% were unique to analysis conducted using the R platform (Table 15). Highlighted

Statistical Approach	Genes
	Methylation Gain
GenomeStudio	<mark>CASD1, HLA-DRB6, LGALS8, SPTBN4, TUBAL3</mark> , ANAPC2, NAT14, OCA2, SYT8, TNNT3, ZNF628
R	AHRR, BAZ2B, C1orf192, CLECL1, HRNBP3, RASA3, RBMY1F, RBMY2FP, SPINK5, TIAL1, TTC22, TUBB8
Both Approaches	ATP8B3, BTNL9, C21orf29, C3orf50, CCS, CNST, COPB1, DCAF11, INSC, KRTAP12-3, KRTAP12-4, LMX1B, MAGI2, MIB2, MRI1, NFIC, PIWIL2, RELN, RHPN1, SNX26, STAG3L4, TAGLN3, TAS2R60, TP73, UST
	Methylation Loss
GenomeStudio	<mark>CLPTM1, FAM120B, FAM69B, KIAA1199, MTUS2, TIMP2,</mark> <mark>VIPR2, ZNF137</mark> , BMP8B, FLJ37201, FLJ43860, FOXK2, JRK, KCNH6, MRGPRX2, PPIE, SND1, SULF2, ZNF490, ZNF665
R	ADORA3, BRMS1, CCDC50, MYO10, NCRNA00052, OPRM1, RIN1, TTTY12, TTTY18
Both Approaches	ACTN3, AKR1C2, ANXA2, ARID1B, B4GALNT3, C2orf69, CAMK1D, COL11A2, CUL3, DEFB128, DMBX1, FAM124B,FRMD4A, GAP43,GCK, JPH3, KCNK7, KRTAP27- 1, LAMB1, MAPK10, MGMT, MORN1, PRKCA, PTPRN2, RAB11B, SAMD4A, SERPINF2, SLC39A14, TOP1MT, WDR36, ZMAT2

Table 15. Differentially Methylated Genes by Statistical Approach

genes in the table are associated with SNPs and had been excluded from analysis in the R environment because SNPs disturb the accuracy of the Infiniuim probes as described in the methods section (Bibikova et al., 2011b; Dedeurwaerder et al., 2011).

#### Validation of Methylation

In order to validate the methylation identified by the Illuminia Infinium 450K array, 6 CpG sites (3 = gain of methylation; 3 = loss of methylation) identified as differentially methylated by GenomeStudio analysis were selected. Sites that were at the extremes of differential methylation, associated with known genes and that primers could be developed were used (Table 16). Six participants were selected at random (3 GBS positive; 3 GBS negative) to validate the array. Participant samples that were sent immediately for sequencing after bisulfite treatment were too poor of quality

UCSCREFGENE	Sequence
RHPN1fwd	GGATGTATTTTTTTTTTTTTTGGTGG
RHPN1rev	CCTCACCCAAATAAACCCTACT
HLA-DRB6fwd	TATTTTAGGATGGATTAGGAGAAAAA
HLA-DRB6rev	CAAAAATTTATAAACACTTCAACAATAC
MRI1 fwd	AATTTTTGATTTTAAGTGATTTGTT
MRI1rev	AAACTATTTCTAAACCATTTTCTACTC
ANXA2fwd	TTGAGGAAAAATAATAAAGAGTTATTAGAT
ANXA2rev	AACCTAAACAATACCATTCAAAACAA
GAP43fwd	TTTAGGTGTGTGTTTTTTTTTTAGGA
GAP43rev	TAACCTTATCTAATTTATCATTTTAACAAC
CUL3fwd	TAGGGGAAAATTGAGGTTATAAGAAG
CUL3rev	TCCTCCTACAATACTAAAATTACAAAC

#### **Table 16. Primer Sequences**

for sequencing as anticipated (Appendix G). The CpG sites associated with ANXA2 (loss

of methylation in GBS positive women) and RHPN1 (gain of methylation in GBS

positive women) were successfully cloned and were of sufficient quality for sequencing.

The DNA used for primers designed with ANXA2 and RHPN1 CpG sites are in Figure 9

below, as the product for these sites were successfully cloned and sequenced Figure 10.

In Figure 9, CpG sites within the primer products are blue text and the CpGs identified as

differentially methylated by GBS status are highlighted yellow. The gray area the DNA

added to identify primers, as described in the methods chapter, and the primer sequence.

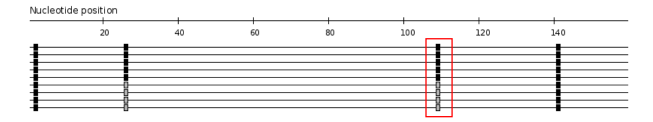
ANXA2>hg19\_dna range=chr15:60643907-60644407 5'pad=250 3'pad=250 strand=+ repeatMasking=none

GGCCACATTCACTTACCCAGGTTCAGGAAAGCATTTTCCAGGTCTCCTTT AACCTCTTTCCTGATGCTTTCCAACATGTCATAAGGGCTGTAACTCTTGT ACCTATCAAATACTGAGGAAAAACAACAACAAAGAGTTATCAGATCCGAGCCA CTAGTCAAAGCTGTCAACGATCACCCACCTAGTTTTATGCACCATAATTT TTTTAAAAATTGAGGATGATCACAGCATCCTAGGAGCTTAGAGGTTACCA CCGTGACCAGAGCCAACATTGGCCAAGTTTGTCGTGGAACAGCCATACCA CCTGTCCTGAATGGCACTGCCCAGGCCACATATTTGGACCATCTCTATCT CCCCTGAGTGGAACCCATTCCATCCGAAAACCATAGGAAACAGTACAGAG CATGCACCAAAGTCCACTACTTCAACAAATAATGGCAAGACCAAATGATC ATCAAACAAGAAGGAGCTGCAGAATAAAGCACCAAATGCAGAAACTATTT G

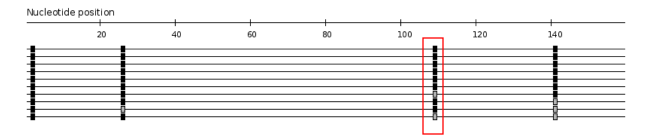
RHPN1>hg19\_dna range=chr8:144457427-144457927 5'pad=250 3'pad=250 strand=+ repeatMasking=none

Figure 9. ANXA2 and RHPN1 DNA, Primer, and Product Sequences

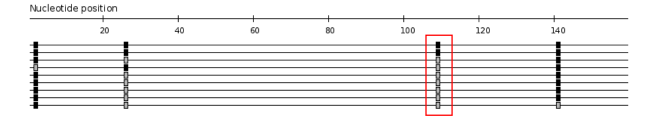
# ANXA2 Participant 10 GBS+ beta value Raw = 0.5709; GS = 0.5638; R= 0.5969



ANXA2 Participant 2 GBS- beta value Raw = 0.8634; GS = 0.8490; R = 0.9388



ANXA2 Participant 14 GBS+ beta value Raw = 0.5409; GS = 0.5344; R = 0.5589



ANXA2 Participant 6 GBS – beta value Raw = 0.8833; GS = 0.8723; R = 9471

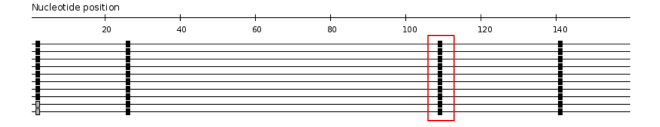
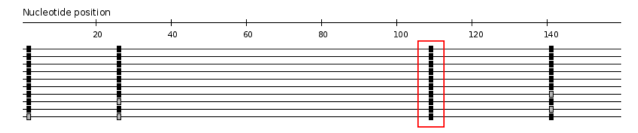
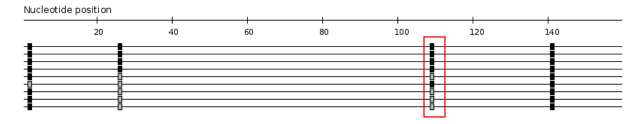


Figure 10. Methylation at ANXA2 and RHPN1 CpG Sites of Cloned Sequences

## ANXA2 Participant 8 GBS- beta value Raw = 0.8901; GS = 0.8768; R = 0.9539



ANXA2 Participant 16 GBS + beta value Raw = 0.4998; GS = 0.4939; R = 0.4997



RHPN1 Participant 10 GBS+ beta value Raw = 0.8229; GS = 0.8186; R = 0.8920 Nucleotide position

	20	40	60	80	100	120	140	160	180	200
		1 1			1 1					
		<u>i i</u>			0					
					1 1					
Ŭ <b>I</b>		<del>i i</del>		ō						

RHPN1 Participant 2 GBS- beta value Raw = 0.4076; GS = 0.4058; R = 0.3845

acieocide po	Sicion									
	20	40	60	80	100	120	140	160	180	200
				-						
					1 1					
•	•••			— i —						— i
		i i								
		1 1						111		
	•••			<b>i</b>	1 1					
•					• •					

RHPN1 Participant 14 GBS+ beta value Raw = 0.8049; GS = 0.8005; R = 0.8855

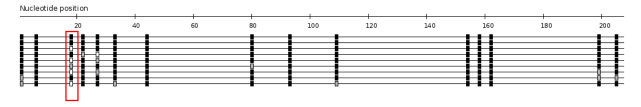
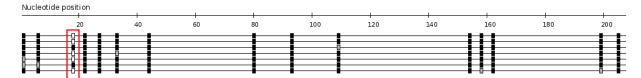
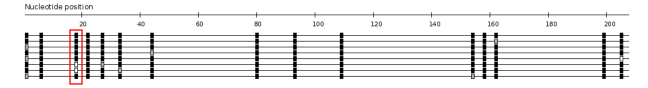


Figure 10. Cont.



RHPN1 Participant 8 GBS- beta value Raw = 0.3858; GS = 0.3842; R = 0.3629

RHPN1 Participant 6 GBS – beta value Raw = 0.3915; GS = 0.4073; R = 0.3684



RHPN1 Participant 16 GBS + beta value Raw = 0.7804; GS = 0.7764; R = 0.8620

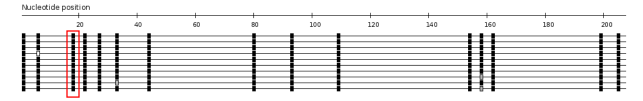


Figure 10. Cont.

In Figure 10, the CpC sites that were identified as having significantly different methylation have a red box around the CpG site in the DNA product sequence. Each figure are cloned sequences for one participant. The notation above the each participant figure indicates the primer by name, participant, GBS status, raw Infinum beta value, and normalized betas calculated by GenomeStudio software and the R programming environment. Black squares indicate the CpG site was methylated, gray squares indicated an unmethylated CpG and white squares indicate it could not be determined. The percent of cloned CpG site methylation was manually calculated by dividing the number of methylated CpGs for the specific site by total number of CpGs that methylation could be determined for the site for each of the participant clones (Table 16). Methylation percent

					ANXA2	KA2			RHPNI	<b>NI</b>	
Participant	Participant Sentrix_ID Position	Position	GBS	Raw beta GS beta	GS beta	R beta	Clones	Clones Rawbeta GS beta	GS beta	R beta	Clones
	5806636022 R01C02	R01C02	Negative	0.5228	0.5133	0.5339		0.8029	0.7975	0.8897	
7	5806636023 R01C02	R01C02	Negative	0.8634	0.8490	0.9388	0.8000	0.4076	0.4058	0.3845	1.0000
С	6229017070 R02C02	R02C02	Negative	0.9026	0.8906	0.9599		0.4405	0.0439	0.4194	
4	6229017070 R04C02	R04C02	Negative	0.9072	0.8966	0.9539		0.1235	0.1231	0.0750	
5	9379082114 R03C01	R03C01	Negative	0.0766	0.0755	0.0473		0.4089	0.4073	0.3906	
9	9379082114 R05C01	R05C01	Negative	0.8833	0.8723	0.9471	1.0000	0.3915	0.4073	0.3684	1.0000
7	9379082114 R01C02	R01C02	Negative	0.5122	0.5034	0.5184		0.0621	0.0619	0.0400	
8	9379082114 R02C02	R02C02	Negative	0.8901	0.8768	0.9539	1.0000	0.3858	0.3842	0.3629	1.0000
6	9379082114 R03C02	R03C02	Negative	0.9273	0.9142	0.9711		0.4361	0.4344	0.4219	
	Average beta GBS		negative	0.7206	0.7102	0.7583		0.3843	0.3406	0.3725	
10	5806636023 R03C01	R03C01	Positive	0.5709	0.5638	0.5969	0.5556	0.8229	0.8186	0.8920	0.9000
11	6229017070 R01C02	R01C02	Positive	0.8768	0.8632	0.9458		0.8318	0.8267	0.9111	
12	9379082114 R01C01	R01C01	Positive	0.0718	0.0708	0.0491		0.7830	0.7779	0.8781	
13	9379082114 R02C01	R02C01	Positive	0.0460	0.0454	0.0244		0.3940	0.3924	0.3727	
14	9379082114 R04C01	R04C01	Positive	0.5409	0.5344	0.5589	0.2222	0.8049	0.8005	0.8855	0.8000
15	9379082114 R06C01	R06C01	Positive	0.0876	0.0862	0.0497		0.8336	0.8299	0.9032	
16	9379082114 R04C02	R04C02	Positive	0.4998	0.4939	0.4997	0.5556	0.7804	0.7764	0.8620	1.0000
17	9379082114 R05C02	R05C02	Positive	0.0985	0.0973	0.0632		0.7999	0.7960	0.8797	
18	9379082114 R06C02	R06C02	Positive	0.5116	0.5053	0.5163		0.8388	0.8352	0.9102	
	Average beta GBS		positive	0.3671	0.3623	0.3671		0.7655	0.7615	0.8327	
	De	Delta beta		-0.3535	-0.3479	-0.3912		0.3812	0.4209	0.4602	

GS = Genome Studio; R = R statistical environment

Table 16. Percent Methylation at Cloned CpG sites

in the cloned sequences was congruent with percent methylation determined by the Illumia Infinium array (Table 16). Participants that had CpG sites where methylation status could not be determined (RHPN1 participant 6, 2 undetermined; RHPN1 Patrticpant 8, 5 undetermined; RHPN1 participant 14, 4 undetermined) did not correlated as strongly as other sites. CpG sites that were identified as significantly different by only one of the software packages had similar beta values, and greater delta beta values did not correlate with significance. Table 17 presents beta results for all participants for one site

	Genome Studio			R environment			
		SND1		RASA3			
Participant	Raw beta	GS beta	R beta	Raw beta	GS beta	R beta	
1	0.4466	0.4435	0.4343	0.3979	0.3958	0.3750	
2	0.0689	0.0683	0.0365	0.8440	0.8376	0.9241	
3	0.8832	0.8766	0.9710	0.4294	0.4273	0.4045	
4	0.8767	0.8711	0.9312	0.4414	0.4397	0.4251	
5	0.4216	0.4193	0.4058	0.8268	0.8219	0.9052	
6	0.8424	0.8372	0.9160	0.4147	0.4130	0.3963	
7	0.8430	0.8358	0.9268	0.4256	0.4233	0.4131	
8	0.8263	0.8196	0.9055	0.8222	0.8165	0.9019	
9	0.8606	0.8548	0.9252	0.8553	0.8501	0.9209	
Average Negaive	0.6744	0.6696	0.7169	0.6064	0.6028	0.6296	
10	0.4428	0.4405	0.4270	0.8536	0.8485	0.9186	
11	0.0519	0.0516	0.0320	0.8801	0.8735	0.9480	
12	0.8383	0.8312	0.9263	0.8177	0.8119	0.9097	
13	0.0532	0.0529	0.0293	0.8271	0.8216	0.9060	
14	0.4477	0.4453	0.4365	0.8244	0.8197	0.9031	
15	0.8578	0.8527	0.9235	0.8430	0.8388	0.9113	
16	0.0532	0.0529	0.0293	0.8362	0.8315	0.9134	
17	0.8666	0.8608	0.9362	0.8343	0.8299	0.9106	
18	0.0688	0.0685	0.0373	0.4285	0.4269	0.4111	
<b>Average Positive</b>	0.4089	0.4063	0.4197	0.7939	0.7891	0.8591	
Delta beta	-0.2655	-0.2633	-0.2972	0.1878	0.1863	0.2295	

Table 17. Beta	Value Simil	arity of Sig	nificant Results

GS= GenomeStudio; R = R statistical environment

Identified as significant using GenomeStudio software (SND1) or the R environment (RASA3). The delta beta for both sites is higher after the R normalization, however only the RASA3 site was identified using the R analysis pipeline as significantly different and SND1 was not. The SND1 site was identified as being significantly different by Genome studio, and RASA3 was not. This type of incongruence is similar at all discordant significant results.

#### Functional Analysis of Methylation

# Research Question 3: Is there a relationship between differentially methylated genes and immune function in pregnant women colonized with GBS?

Since no CpG sites were identified as differentially methylated when FDR is applied to determine significance, results produced using the less stringent alpha of 0.05 for significance were used for functional analysis. Cluster analysis using DAVID revealed differential methylation in women with GBS is primarily related to basic cellular processes. DAVID analysis was performed as described in the methods section independently for DNA methylation results from GenomeStudio (Table 16), R (Table 17), and genes common to both analytical approaches (Table 18). However, none of the functional clusters were significant using FDR of 0.05 and the cell morphogenesis cluster was the only functional cluster that was significant at an alpha level of 0.05 for data output from GenomeStudio, R and for common genes identified by both approaches. The results of the functional analysis using the data generated with this sample yielded different functional clusters from the preliminary data analysis used when designing and conceptualizing this study that clustered more specifically to immune function such as major histocompatibility complex and antigen presentation (Appendix H).

Functional Cluster	Enrichment Score	Genes	p- value	FDR
Keratin	1.26	Methylation Gain KRTAP12-3, KRTAP12-4, ZNF628	0.04	0.98
Cellular metabolic and biosynthetic processes	1.15	LMX1B, ANAPC2, PIWIL2, NFIC, TP73	0.04	0.97
Protein kinase and phosphorylation	0.87	ANAPC2, RELN, TP73	0.10	0.99
Regulation of transcription	0.67	LMX1B, NFIC, TP73	0.17	1.00
Cell cycle	0.64	ANAPC2, PIWIL2, TP73	0.17	1.00
Cytoplasmic membrane- bounded vesicle	0.53	ATP8B3, COPB1, SYT8	0.26	1.00
Transcription factor activity	0.42	LMX1B, NAT14, NFIC, TP73, ZNF628	0.22	1.00
Ion binding	0.31	ATP8B3, LMX1B, CCS, MIB2, RELN, TP73, ZNF628	0.40	1.00
Membrane	0.25	ATP8B3, CASD1, NAT14, BTNL9, CNST, COPB1, PIWIL2, MAGI2, OCA2,	0.53	1.00
		Methylation loss		
Extracellular matrix	1.58	TIMP2, ANXA2, COL11A2, LAMB1	0.02	0.96
Cell morphogenesis	0.84	CUL3, GAP43, LAMB1, PRKCA	0.03	1.00
Regulation of apoptosis	0.75	MGMT, ACTN3, CAMK1D, CUL3, PRKCA	0.18	1.00

# Table 16. Biological Pathways Associated with GBS (GenomeStudio data)

Table 16. Cont.

Functional	Enrichment		р-	
Cluster	Score	Genes	value	FDR
Cytoplasmic membrane- bounded vesicle	0.57	RAB11B, ANXA2, SERPINF2, SND1	0.22	0.98
Phosphorus metabolic process	0.56	CAMK1D, GCK, MAPK10, PRKCA, PTPRN2	0.27	1.00
Protein dimerization activity	0.56	MTUS2, ACTN3, COL11A2, DMBX1	0.18	1.00
Ion transport	0.53	JPH3, KCNK7, KCNH6, SLC39A14	0.13	1.00
Cellular homeostasis	0.46	COL11A2, GCK, JPH3, PRKCA	0.28	1.00
Transcription regulation	0.39	ARID1B, DMBX1, FAM120B, FOXK2, SND1, ZNF490, ZNF665	0.36	0.98
Ion binding	0.34	MGMT, ACTN3, ANXA2, CAMK1D, FOXK2, KCNK7, KCNH6, PRKCA, SLC39A14, SULF2, ZNF490, ZNF665, ZMAT2	0.44	1.00
Nucleotide binding	0.33	RAB11B, CAMK1D, GCK, MAPK10, PPIE, PRKCA, TOP1MT	0.58	1.00
Regulation metabolic and biosynthetic processes	0.28	ARID1B, GCK, PRKCA, SAMD4A	0.42	1.00
Biological adhesion	0.25	ACTN3, COL11A2, LAMB1	0.52	1.00
Zinc ion binding	0.18	MGMT, TIMP2, PRKCA, SLC39A14, ZNF490, ZNF665, ZMAT2	0.43	0.97

# Table 16. Cont.

Functional	Enrichment		р-	
Cluster	Score	Genes	value	FDR
Membrane- enclosed lumen	0.16	ARID1B, MGMT, DMBX1, GCK, SERPINF2, TOP1MT	0.65	1.00
Zinc finger, C2H2- like	0.14	ZNF490, ZNF665, ZMAT2	0.62	1.00
Transmembrane	0.07	MRGPRX2, B4GALNT3, CLPTM1, FAM69B, JPH3, KCNK7, KCNH6, PTPRN2, SLC39A14, VIPR2	0.87	1.00

Functional	Enrichment		р-	
Cluster	Score	Genes	value	FDR
		Methylation Gain		
RNA recognition motif, RNP-1	1.19	RBMY1F, TIAL1, HRNBP3	0.06	1.00
Cellular metabolic and biosynthetic processes	0.97	LMX1B, PIWIL2, NFIC, TP73	0.08	1.00
Induction of apoptosis	0.71	TIAL1, AHRR, TP73	0.09	1.00
Metal-binding	0.65	ATP8B3, LMX1B, RASA3, BAZ2B, CCS, MIB2, RELN, TP73	0.17	1.00
Regulation of transcription	0.64	LMX1B, TIAL1, AHRR, BAZ2B, NFIC, TP73	0.11	1.00
Cytoplasmic membrane- bounded vesicle	0.61	ATP8B3, COPB1, SPINK5	0.21	1.00
Transcription factor activity	0.60	LMX1B, NFIC, TP73	0.18	0.99
Ion binding	0.44	ATP8B3, LMX1B, RASA3, AHRR, BAZ2B, CCS, MIB2, RELN, TP73	0.43	1.00

## Table 17. Cont.

Functional	Enrichment		р-	
Cluster	Score	Genes	value	FDR
Transcription regulation	0.29	LMX1B, TIAL1, AHRR, BAZ2B, NFIC, TP73	0.42	1.00
Intrinsic to membrane	0.06	ATP8B3, CLECL1, RASA3, BTNL9, CNST, PIWIL2, TAS2R60, UST	0.83	1.00
		Methylation loss		
Cell morphogenesis	1.63	CUL3, GAP43, LAMB1, PRKCA	0.01	1.00
Neuron development	0.85	GAP43, LAMB1, PRKCA	0.07	1.00
Phosphorus metabolic process	0.84	CAMK1D, GCK, MAPK10, PRKCA, PTPRN2	0.11	1.00
Nucleotide binding	0.73	RAB11B, CAMK1D, GCK, MAPK10, MYO10, PRKCA, TOP1MT	0.10	0.79
Serine/threonine protein kinase	0.68	CAMK1D, MAPK10,PRKCA	0.09	1.00
Cellular homeostasis	0.67	GCK, JPH3, PRKCA	0.16	1.00
G-protein coupled receptor protein signaling pathway	0.64	ADORA3, GAP43, OPRM1	0.65	1.00
Ion transport	0.52	JPH3, KCNK7, SLC39A14	0.22	1.00
Regulation biosynthetic processes	0.42	ARID1B, GCK, SAMD4A	0.36	1.00
Biological adhesion	0.41	ACTN3, COL11A2, LAMB1	0.39	1.00
Cytoplasmic membrane- bounded vesicle	0.36	RAB11B, ANXA2, SERPINF2	0.39	0.99

## Table 17. Cont.

Functional	Enrichment		р-	
Cluster	Score	Genes	value	FDR
Membrane-	0.28	ARID1B, MGMT, DMBX1, GCK,	0.46	0.98
enclosed lumen		SERPINF2, TOP1MT		
Receptor	0.18	ADORA3, OPRM1, PTPRN2	0.82	1.00
Ion binding	0.09	MGMT, ACTN3, ANXA2, CAMK1D, KCNK7, PRKCA, SLC39A14, ZMAT2	0.81	1.00
Zinc ion binding	0.06	MGMT, PRKCA, SLC39A14, ZMAT2	0.81	1.00
Integral to membrane	0.06	ADORA3, B4GALNT3, JPH3, NCRNA00052, OPRM1, KCNK7, PTPRN2, SLC39A14	0.99	1.00

# Table 18. Biological Pathways Common between R and GenomeStudio

Functional Cluster	Enrichment Score	Genes	p- value	FDR
		Methylation Gain		
Cellular metabolic and biosynthetic processes	1.46	LMX1B, PIWIL2, NFIC, TP73	0.03	1.00
Metal-binding	0.51	ATP8B3, LMX1B, CCS, MIB2, RELN, TP73	0.28	0.99
Regulation of transcription	0.50	LMX1B, NFIC, TP73	0.08	0.99
Intrinsic to membrane	0.06	ATP8B3, BTNL9, CNST, PIWIL2, TAS2R60, UST	0.74	1.00
		Methylation loss		
Cell morphogenesis	1.77	CUL3, GAP43, LAMB1, PRKCA	0.01	0.97
Regulation of apoptosis	1.34	MGMT, ACTN3, CAMK1D, CUL3, PRKCA	0.05	0.98
Extracellular matrix	1.14	ANXA2, COL11A2, LAMB1	0.05	0.98

## Table 18. Cont.

Functional	Enrichment		р-	
Cluster	Score	Genes	value	FDR
Neuron development	0.94	GAP43, LAMB1, PRKCA	0.07	1.00
Serine/threonine protein kinase	0.79	CAMK1D, MAPK10,PRKCA	0.07	1.00
Cellular homeostasis	0.75	GCK, JPH3, PRKCA	0.14	0.99
Phosphorus metabolic process	0.74	RAB11B, CAMK1D, GCK, MAPK10, PRKCA, PTPRN2	0.12	0.83
Metal ion transport	0.59	JPH3, KCNK7, SLC39A14	0.19	0.99
Cellular biosynthetic processes	0.49	ARID1B, GCK, SAMD4A	0.31	1.00
Biological adhesion	0.49	ACTN3, COL11A2, LAMB1	0.31	0.98
Cytoplasmic membrane- bounded vesicle	0.46	RAB11B, ANXA2, SERPINF2	0.31	0.98
Membrane- enclosed lumen	0.43	ARID1B, MGMT, DMBX1, GCK, SERPINF2, TOP1MT	0.31	0.99
Ion binding	0.19	MGMT, ACTN3, ANXA2, CAMK1D, KCNK7, PRKCA, SLC39A14, ZMAT2	0.64	1.00
Zinc ion binding	0.10	MGMT, PRKCA, SLC39A14, ZMAT2	0.68	1.00
Integral to membrane	0.02	B4GALNT3, JPH3, KCNK7, PTPRN2, SLC39A14	0.80	1.00

## Summary of Results

In summary, the results from this study indicate that there are no significant differences in DNA methylation between women with and without GBS colonization when FDR is used to determine significance of DNA methylation differences that are greater than 20% between groups. However, if a less stringent p-value of 0.05 is used there are a small number of CpG sites that have significant differences with greater than a 20% difference in methylation. The number of significantly different CpG sites identified using different software for analysis varies (141 versus 125). No significant association was found between serum TNF- $\alpha$ , IL-6, IL-10 or vitamin D (25[OH]D) levels and maternal GBS colonization status. Lastly, analysis of functional pathways did not find a correlation between differentially methylated genes and genes directly related to cytokine production or specific immune pathways. No functional clusters were significant when applying FDR for significance; and only the cell morphology functional cluster was significant for output from both GenomeStudio and R data when an alpha of 0.05 was applied.

#### CHAPTER V

#### DISCUSSION

GBS sepsis continues to be the leading cause of infectious neonatal morbidity and mortality despite current practice guidelines to prevent the transmission of GBS from mothers to their infants (Phares et al., 2008). The primary purpose of this exploratory study was to identify variants in maternal blood that are associated with maternal GBS colonization in order to assist with the development of more accurate screening tools and/or assist in identifying to targets to prevent maternal GBS colonization. It is unknown why GBS selectively colonizes one third of pregnant women, placing the health of women and their offspring at risk. Currently in the US, all pregnant women are screened for GBS colonization between 35-37 weeks gestation (Verani et al., 2010). However, significant false negative screening results and infant illness despite maternal antibiotic treatment during labor requires further investigation to identify a biological reasons as to why certain women are preferentially colonized with GBS (Lin et al., 2011; Towers et al., 2010). For this study, variants that can be measure in the serum that are increasingly used for monitoring and diagnosing other clinical conditions, were investigated to determine if there was any association between serum levels and maternal GBS colonization status. This contribution to science is significant because the relationship between DNA methylation patterns, TNF- $\alpha$ , IL-6, IL-10, and vitamin D (25[OH]D) with maternal GBS colonization status have not previously been reported.

#### Maternal Serum Cytokines and Vitamin D (25[OH]D)

Research Question 1: Are serum levels of TNF- $\alpha$ , IL-6, IL-10, and vitamin D (25[OH]D) different in pregnant women with GBS colonization than pregnant women without GBS

### colonization?

This study was designed to capitalize on data that was collected for a previous study. Other clinical indicators could potentially be used to identify pregnant women at increased risk for GBS colonization that have not previously been considered. Clinical indicators, such as serum cytokine levels that reflect immune system functioning, could also be associated with altered DNA methylation patterns. Variability in serum markers associated with immune function and vitamin D (25[OH]D) have been previously reported and utilized as a prognostic indicator of disease states, such as respiratory infections (Chesney, 2010), human immunodeficiency virus infections (Fahey et al., 1990), pancreatitis (Pezzilli et al., 1995), and depression (Kiecolt-Glaser & Glaser, 2002). Research investigating the role of vitamin D (25[OH]D) as an immune function modulator has increased dramatically in recent years (Figure 3) and could offer a cost effective intervention target if low serum vitamin D (25[OH]D) levels are associated with GBS colonization.

Multiple alterations in immune function are necessary during pregnancy to prevent the mother's body from perceiving the developing fetus as a foreign pathogen. Serum markers of immune function are being increasingly evaluated in pregnancy because alterations occur throughout normal pregnancy and inappropriate levels contribute to the development of pathology during pregnancy (Ponsonby et al., 2010). In a recent study, Fichorova et al. (2011) identified patterns in immune function markers

that were specific to the type of bacteria present in the vaginal mucosa and placenta of pregnant women. Notably, they found TNF- $\alpha$ , IL-1 $\beta$ , IL-6, IL-8 and ICAM-1 were elevated when pathogenic organisms associated with bacterial vaginosis were present. Furthermore, *Lactobacillus*, which colonize the vaginal mucosa and are not pathogenic, suppress pathogenic strains and downregulate pro-inflammatory cytokines (Donato et al., 2010; Othman et al., 2007; Zeuthen et al., 2010). However, no evidence could be found in the literature to suggest that patterns of immune function serum markers have been explored with regards to GBS colonization or infection. Additionally vitamin D (25[OH]D), a known modulator of immune function and deficiencies, has been associated with susceptibility to infectious diseases (Chesney, 2010). It is generally accepted that serum vitamin D (25[OH]D) levels must be above 20 ng/ml during pregnancy to maintain normal physiologic processes and fetal development. Recently, experts have noted vitamin D (25[OH]D) levels greater than 32 ng/ml are necessary to support all physiologic processes (e.g. infection prevention) that require vitamin D (25[OH]D) for optimal functioning (ACOG, 2011; Holick et. al., 2011). For this study, TNF- $\alpha$ , IL-6, IL-10 and vitamin D (25[OH]D) were selected specifically for analysis because of the reported association with infectious diseases and action during pregnancy. The serum cytokine and vitamin D (25[OH]D) levels were evaluated throughout the pregnancy in order to identify any difference in how the levels change throughout pregnancy, as well as independently during each trimester. Laboratory results were also evaluated to identify any direct correlation between serum cytokine and vitamin D (25[OH]D) levels.

First, correlation tests indicated that vitamin D (25[OH]D) levels were not correlated with serum TNF- $\alpha$ , IL-6, or IL-10 levels during pregnancy. The initial hypothesis of the study was that vitamin D (25[OH]D) would be correlated with TNF- $\alpha$ , IL-6, or IL-10 levels because vitamin D (25[OH]D) modulates immune function. Because there was no correlation between the cytokines and vitamin D (25[OH]D); vitamin D (25[OH]D) was not used as a covariate and repeated measures ANOVA was completed using serum vitamin D (25[OH]D), TNF-α, IL-6 and IL-10 independently instead of the originally planned repeated measures ANCOVA (with vitamin D (25[OH]D) as the covariate). In addition to there being no correlation between serum vitamin D (25[OH]D) levels and the serum cytokines, no significant association between serum TNF- $\alpha$  or IL-6 levels and maternal GBS colonization status was identified. In vitro experiments have indicated there is an increase in TNF- $\alpha$  production in neonatal and adult peripheral mononuclear cells (Berner et al., 2002) and human epithelial cells (Mikamo et al., 2004) exposed to GBS. Additionally, TNF- $\alpha$  and IL-6 levels increase systemically in murine models when the mice are inoculated with GBS resulting in high mortality rates (Puliti et al., 2000). Because TNF- $\alpha$  levels increase in laboratory and animal studies as a result of GBS exposure, this investigator hypothesized that TNF- $\alpha$  levels may be elevated in pregnant women who are GBS positive. In this study, TNF- $\alpha$  levels were only elevated in pregnant women with GBS during the third trimester and the difference was not statistically significant. Furthermore, results from the one-way MANOVA indicated TNF- $\alpha$  has no effect (partial  $\eta^2 < 0.001$ ) on GBS colonization status in pregnant women. Given this very low  $\eta^2$ , it is unlikely that any significant difference would be identified in

TNF- $\alpha$  values in pregnant women with and without GBS colonization; even with a larger sample size.

There is an increased IL-6 production in vitro, and in animal models, with exposure to GBS (Berner et al., 2002; Mikamo et al., 2004; Puliti et al., 2002). The increase in IL-6 in response to GBS exposure has not been verified or validated in human studies. This study evaluated the level of serum IL-6 to determine if a similar increase in IL-6 production occurs in pregnant women in response to GBS colonization and found no statically significant effect. Results from the one-way MANOVA indicated IL-6 has no effect (partial  $\eta^2 < 0.001$ ) on GBS colonization status in pregnant women. It is unlikely that even with a larger sample size any significant difference would be identified in IL-6 values in pregnant women with and without GBS colonization.

Two previous studies in murine models identified an increase in IL-10 levels related to GBS exposure or infection (Bebien et al., 2012; Madureira et al., 2011). No statistically significant elevation in IL-10 levels in pregnant women who were GBS positive were observed at any point during pregnancy. Because of the small sample size, we calculated the mean serum level values and conduct a one-way MANOVA to correct for confounding resulting from data being collected over time, determine effect size and observed power. Because we found a negligible effect (partial  $\eta^2 = 0.006$ ) of IL-10 on GBS colonization status, it may be warranted to repeat the study with a larger sample size.

Serum vitamin D (25[OH]D) levels had a small effect on GBS colonization status. The normal increase in serum vitamin D (25[OH]D) levels during pregnancy that may be involved in preventing infection and colonization with pathogenic bacteria during normal

pregnancy was not evident in this study (De-Regil Luz et al., 2012). Surprisingly, multivariate analysis revealed that GBS positive women are more likely to have higher serum vitamin D (25[OH]D) levels. However, both GBS positive (29.16 ng/ml) and GBS negative women (24.59 ng/ml) had serum vitamin D (25[OH]D) levels below the 32 ng/ml recommended for optimal physiologic functioning during pregnancy. Both groups of women likely had low vitamin D (25[OH]D) levels because of the fact that the primary study was undertaken in the high northern latitude. Unfortunately, previous studies evaluating vitamin D (25[OH]D) levels related to GBS colonization status could not be found in the literature. Therefore, it is unclear how these results should be interpreted. A meta-analysis conducted by Thorne-Lyman and Fawzi (2012) revealed it is unknown how vitamin D (25[OH]D) relates to maternal infections since the relationship between vitamin D (25[OH]D) and immunity has only recently been established. Vitamin D (25[OH]D) levels across pregnancy will be evaluated in a larger cohort of women in an upcoming follow-up study to determine if the same results persist. Repeating the analysis with a larger cohort may result in better understanding of the significance of these study results and how vitamin D (25[OH]D) is related to GBS status during pregnancy. Posthoc power analysis indicated that replicating the study with a larger sample size may yield significant differences in vitamin D (25[OH]D) levels associated with GBS colonization status because serum vitamin D (25[OH]D) levels appear to have some effect on GBS colonization status (partial  $\eta^2 = 0.068$ ).

#### Maternal DNA Methylation

Research Question 2: Are DNA methylation patterns different between pregnant women with GBS colonization and those without GBS colonization?

DNA methylation is currently the most well understood epigenetic modification and is increasingly being integrated into clinical nursing research. In this study, the investigator used previously collected peripheral blood samples in order to complete the study in a timely and cost effective manner. The exploratory nature of this genome wide DNA methylation analysis allowed the investigator to quantify methylation of individual CpG sites. Specifically, DNA methylation in peripheral white blood cells that were collected during the first trimester of pregnancy in women colonized with GBS in late pregnancy were compared to women who screened negative for GBS in the third trimester of pregnancy. Preliminary analysis of a subset of individuals by the investigator (n=6), initially indicated that over 1,000 potential early pregnancy DNA methylation differences existed between women with and without late pregnancy GBS colonization. Since DNA methylation is an epigenetic modification that can result in altered gene expression, with the potential to impact health and disease susceptibility, this study was designed to see if the preliminary differences identified in a very small sample (2 GBS) positive, 4 GBS negative) would persist in a larger sample. Previous studies have identified potentially useful, clinically relevant DNA methylation biomarkers for preeclampsia using a sample size of n=6/group (Anderson et al., 2013). For this study, the sample size was increased to n=9/group, which reduced the number of statistically significant differentially methylated sites from over 1,000 genes between women with and without GBS colonization to 141 CpG sites using the GenomeStudio software. Due to the small number of significant genes and the drastic decrease in potentially different CpG sites, it is premature to assume these sites may be an early biomarker for GBS colonization. Support for this assessment will be evident in the discussion of the variation

in the number and actual genes associated with differentially methylated CpG sites using different software for statistical analysis. Prior to this study, no other papers could be identified in the literature investigating host DNA methylation patterns related to colonization with a certain microbe. As discussed in chapter 2, this is an area of inquiry that will likely increase over time because of the role that methylation plays with cell differentiation and memory in immune cells (Bobetsis et al., 2007; Torsten Olszak et al., 2012; Schaub et al., 2009; Tolg et al., 2011).

Of the 18 women included in the methylation analysis for this study, data extracted from the medical record indicated no difference in baseline characteristics between GBS positive and GBS negative women. Further, no significant differences in maternal co-morbidities that may indicate, or cause, altered immune function; such as asthma, infections or antimicrobial usage were found. Because some of the normalizations features incorporated into the analytic component of GenomeStudio software are proprietary, it has limited functionality in assessing and presenting differentially methylated data and may limit reproducibility (Gentleman et al., 2004; Smyth, 2005). Data analysis pipelines for biology and bioinformatics were design because "the primary motivations for an open-source computing environment for statistical genomics are transparency, pursuit of reproducibility and efficiency of development" (Gentleman et al., 2004, p. R80.2). Therefore, statistical analyses were also completed using the R statistical environment to ensure accurate normalization and interpretation and reproducibility of analysis of the raw data. Additionally, Hansen et al. (2013) noted that it was unclear as to what process is used for normalization by GenomeStudio because it is not explicitly publicized. Due to the lack of transparency of

normalization methods used in GenomeStudio analytic software that limits reproducibility of analysis, biostatisticians and R developers recommend not using GenomeStudio to conduct statistical analysis. However, R and GenomeStudio are both used extensively to conduct methylation analysis. Because both programs are frequently used, the results produced using GenomeStudio and R to conduct statistical analysis were included in the results section. Some of the differences in the results are likely a direct result of different normalization methods of the raw data in each platform. Each result can be reproduced using the specified statistical software, although results differ between software packages.

For the actual statistical analysis of DNA methylation using the R statistical environment, M- values were used, instead of beta values because M-values are homoscedastic across different levels of methylation. Given that beta values exhibit severe heteroscedasticity at the methylation extremes, M-values are the superior choice for conducting valid differential and statistical analysis (Du, Kibbe, & Lin, 2008). However, M-values are difficult to interpret clinically for relevance because the transformed negative values are not biologically interpretable. Therefore, after using the logit transformed values (M values) for statistical analysis, the data were reported using the original untransformed beta values to allow for clinical interpretation of the findings. In fact, beta values are most frequently reported because beta values represent a percentage of methylation, and therefore are more biologically meaningful (Du et al., 2010).

Regardless of the statistical platform used for the data analysis, we did find over 100 CpG sites with significantly different methylation between women who were GBS

positive or GBS negative when applying an alpha of 0.05. Of the sites identified by both GenomeStudio and R statistical environment, there were 56 common genes. However, it is expected that statistically by chance 24,250 CpG sites would be identified as differentially methylated when using an alpha of 0.05 (485,000 sites in the array \* 0.05 = 24,250). In other words, basic statistics indicate an arbitrary investigator is more likely to find a significantly different methylated CpG site between GBS positive and negative women, than the sites identified using an alpha of 0.05 actually being significant. Similar analytic approaches for identifying biomarkers, such as the NIMBL package for Matlab, likely also lack the power to identify sub-sample heterogeneity or reasonably identify biomarkers in a sample of this size when not using FDR to determine significance. Therefore, conducting additional testing assessing for significant in this sample would not yield valuable biomarker information because no sites are significantly different when FDR is applied to determine significance. It is imperative to generate empirical estimates of test statistics (and p-values) via bootstrapping methods for small sample sizes, coupled with FDR to appropriately safeguard against over-interpretation of microarray data (Benjamini & Hochberg, 1995; Tusher, Tibshirani, & Chu, 2001; York, 2003). Furthermore, recommendations for design and analysis of epigenome-wide association studies, such as this one, include using multiple-testing adjustments and validating the methylation in a similar but different cohort using a different laboratory methodology (Michels et al., 2013). Analysis of methylation on this study cohort using FDR indicate no CpG sites have significantly different methylation using any FDR cutoff value (adjusted p value= 0.99992 for all values, Appendix I). The lack of significance maybe attributed to small sample size and the use of peripheral blood instead of vaginal

epithelial cells. Michels et al. (2013) stated "As natural variation affects DNA methylation, larger sample sizes will typically be required for EWAS than for GWAS for any given phenotype, even when the most technically sophisticated assays are used" (p. 952), therefore it is premature to assume that any of the CpG sites identified in this study with 18 participants will yield reliable biomarker results regardless of approach. In future studies, the same analysis will be completed using a larger sample size of GBS positive and GBS negative women who have already had genome-wide methylation analysis completed. This investigator suspects that repeating the analysis on a larger cohort of women may further decrease the number of statistically significant differentially methylated sites identified, or result in completely different findings between the two groups of women. Such replication will also enable the investigator to determine if genes identified by only one of the approaches (GenomeStudio or R) are no longer significant. Further analysis with a larger sample or statistical simulations, such as bootstrapping that are beyond the scope of this dissertation, may be able to better identify why the results differ when using different software to perform statistical analysis.

Additional studies will also be needed to determine the significance of DNA methylation on gene expression. RNA was not collected in the parent study, and therefore gene expression studies using RNA could not be completed for this study. However, based on the location of differential DNA methylation, it is possible that the differentially methylated CpG sites associated with genes may have an effect on gene expression. CpG islands, dense regions of cytosine and guanine dinucleotides, contribute to the regulation of gene transcription and subsequent gene expression (Deaton & Bird, 2011). Approximately 72% of known gene promoter regions are associated with CpG islands

(Saxonov, Berg, & Brutlag, 2006). Results from the analysis conducted in the R environment identified 15 CpG sites that were within CpG islands and 44 CpG sites were located in the regions flanking the islands that may also result in altered gene expression patterns (Doi et al., 2009). Upon follow-up studies methylation at these sites will be assessed, since they are most likely related to gene expression. Appropriate samples will also be collected to perform RNA and protein analysis in the next study.

If a selective and specific biomarker panel for GBS colonization based on differential methylation patterns can be developed after repeating the analysis with a larger cohort, it could be useful for identifying women at risk for poor pregnancy outcomes (e.g. miscarriage, preterm birth, premature rupture of membranes and maternal infections) that occur as a result of GBS prior to 35-37 weeks of pregnancy. Since DNA methylation vitally contributes to programming memory in immune cells, altered methylation patterns in women with GBS could represent a novel target for designing novel treatment and prevention modalities.

Biologic Functions Associated with Altered Methylation Research Question 3: Is there a relationship between differentially methylated genes and immune function in pregnant women colonized with GBS?

Although no CpG sites were differentially methylated when applying FDR for significance, functional analysis was conducted using DAVID bioinformatics data base to determine if the differentially methylated genes identified using an alpha of 0.05 may be related to immune function or inflammation. A recently published manuscript by Laayouni et al. (2014), identified alterations in 20 genes associated with immune function in populations exposed to *Yersinia pestis* that have persisted over time and resulted in

altered predisposition for autoimmune disorders in individual of European descent. Laayouni's team found variant SNPs in genes that alter how the immune system responds to Y. pestis. They found that the production of pro-inflammatory cytokines is increased in response to Y. pestis. The increased inflammatory response allowed for some of the population to be resistant to, or heal from, the Black Plague. These SNP variations are not seen in populations that were out of the endemic area during the Black Plague. Laayouni's group suggested that the SNP variations, driven by pathogenic exposure, contribute to the increased prevalence of autoimmune disorders in populations of European descent where Black Plague was endemic. The SNP variants are not present in populations not exposed to Y. pestis and the populations also exhibit lower prevalence of autoimmune disorders. Lauyouni's study further supports that genetic variation can be driven by, and contribute to, pathogen specific immune response that persists for generations. Multiple studies have identified altered DNA methylation patterns that have occurred in response to, or as a result of, exposure to specific pathogens (Bobetsis et al., 2007; Mikovits et al., 1998; Tolg et al., 2011). This study was the first to evaluate DNA methylation patterns in women with GBS colonization and functional analysis reveals these changes may play contribute to colonization susceptibility.

Genes incorporated into significant functional clusters using the DAVID bioinformatics software were independently searched within the GeneCards® database, The Human Gene Compendium Encyclopedia (http://www.genecards.org/). Functional cluster analysis was completed using DAVID bioinformatics software. None of the functional categories were significant when FDR is applied to determine significance of the clusters identified. However, for genes identified by using both GenomeStudio and R

software as being differentially methylated, the cell morphogenesis functional cluster has potential to offer mechanistic insight into GBS colonization and was significant using and alpha of 0.05 (p=0.01). The functional cluster was associated with methylation loss in GBS positive women. The cell morphogenesis cluster includes four differentially methylated genes and has an enrichment score of 1.77, which is above the suggested 1.3 cutoff value indicating likely involvement in biological process (Huang et al., 2009a). What makes this particular cluster interesting is that the genes within the cluster are associated with various immune functions and pathways. Gómez et al., (2010) determined that there is an association with PRKCA and bacterial vaginosis. This gene is associated with abnormal bacteria in the vagina and a similar association may exist with GBS susceptibility. The CUL3 gene is in a SuperPaths specifically related to antigen processing and the adaptive immune system (Andérica-Romero, González-Herrera, Santamaría, & Pedraza-Chaverri, 2013; Pintard, Willems, & Peter, 2004; Singer, Gurian-West, Clurman, & Roberts, 1999). It is possible that the loss in methylation in women with GBS alters antigen presentation and how the body responds to GBS (e.g. allowing colonization or clearing the bacteria). The GAP43 gene has been associated with inflammatory disease processes including contact dermatitis (El-Nour et al., 2006) and cutaneous malignant melanoma (Reed, Finnerty, & Albino, 1999). Since the gene is hypomethylated in women with GBS, it is possible there is increased cutaneous inflammation which is damaging to normal flora and creates a niche for GBS to colonize. Additional research investigating expression levels, protein products, and associated clinical outcomes, could be beneficial for the genes in this functional cluster.

There were two additional functional clusters with enrichment scores above the 1.3 threshold for biological significance: cellular metabolic and biosynthetic processes (1.46, p=0.03) associated with genes that were hypermethylation and regulation of apoptosis (1.34, p = 0.05) associated with genes that were hypomethylated. Upon searching in GeneCards and PubMed, there does not appear to be any literature directly associating the genes examined in this study to with immune variations related to infections in either of the aforementioned pathways. There are 1,166 publications associating the genes identified with various cancers and neurological ailments, but none of the studies directly pertaining to infectious disease processes. Therefore, it seems unlikely that the genes in this cluster will offer any mechanistic insight unless there are indirect linkages to inflammatory processes. However, genes identified in this cluster may still be useful as clinical biomarkers in the future for identifying carriers, or women at risk for colonization in early pregnancy to prevent preterm labor or other poor health outcomes if the results are replicated in a larger cohort.

Two of the genes (CUL3 and PRCKA) in the apoptosis cluster are the same as in the cell morphogenesis cluster. The remaining genes in the apoptosis functional cluster do not appear to have any direct linkages to immune processes related to infection. There were 1,462 publications identified for the three genes that did not overlap with the morphogenesis cluster that were associated primarily with tumors, cancers, and muscular dystrophies. Similar to the metabolic and biosynthetic cluster, it seems unlikely that this cluster will offer any significant mechanistic understanding to GBS colonization. However, these genes may also be useful clinical biomarkers in the future after additional analysis is conducted for identifying GBS carriers or individuals susceptible to GBS

colonization. Further evaluation in a larger cohort and gene expression data may assist in determining if these genes are useful as clinical biomarkers for GBS colonization. Future studies investigating the relationship between PRKCA, CUL3, and GAP43 genes and GBS colonization may offer mechanistic insight and provide targets for future GBS treatment or to develop colonization prevention strategies.

#### Nursing Implications

Advances in epigenomic research are beginning to contribute significantly to scientific understanding of how environmental factors may contribute to various disease processes. This study is the first to assess laboratory values that are increasingly being used for nursing research (cytokines, vitamin D and DNA methylation) related to GBS colonization status in pregnant women. Existing gene-environment interaction models (Figure 1) were adapted (Figure 2) in order to guide the approach for this nursing research study to investigate if quantitative variations in cytokines or DNA methylation levels identified in in pregnant women's could be used to identify women at risk for GBS colonization. Although this study did not identify any significant differences between women colonized with GBS and women without GBS colonization, this study adds a novel model and approach method that can be used in future nursing research that can be modified as research methodology continues to evolve. Expanding on existing research models to bridge the gap for nurses to conduct translational research to improve outcomes will have implications for nursing research, practice, education and policy.

### Nursing Research Implications

Nurses are uniquely poised to accelerate the translational arm of epigenomic research to better assist clinical populations of interest to attain and maintain optimal health functioning. Conducting research that critically examines environmental exposures

and unique epigenomic signatures will allow for the discovery of new treatment targets and the ability to create new disease prevention strategies ranging from diet modification to driving forward policy chance to protect public health and well-being. This is the first study investigating cytokines, vitamin D, or DNA methylation levels in peripheral blood to identify potential associations with GBS colonization susceptibility. Historically, much of the epigenomic literature is dominated by cancer studies. However, studies investigating the relationship between epigenomic alterations and complex diseases, other than cancer, have been increasing in recent years. It is important to note that identifying aberrant DNA methylation patterns alone will not explain why or how it was altered, how to intervene, or help the patients avoid acquiring abnormal patterns.

#### Nursing Practice Implications

A holistic approach for investigating the impact of epigenomic alterations on health status is a necessity if science intends on using epigenomic information to improve health. Other lifestyle patterns (diet, lifestyle, stress) and exposures must be assessed to identify interactions that may be causing the altered DNA methylation pattern. Nursing clinical assessments can help illuminate human-environment interactions, the endogenous and exogenous factors in the model developed to guide this study (Figure 2), that may be altering DNA methylation patterns that cannot be identified by studies using cell or animal models. Studies involving actual clinical populations are needed, specifically for disease processes that are inflammatory in nature because the immune response in murine models does not correlate with the human inflammatory response (Seok et al., 2013). Although it is near impossible to select populations to eliminate all confounding variables, perhaps it is time to embrace studies that acknowledge and address

confounding variables. The environment has an undeniable impact on health and disease states, therefore to completely eliminate all confounding variables from bench studies may explain some of the barriers encountered when translating research from bench to bedside. Humans do not live in a well-controlled, isolated environment; and their environment will alter how they respond to treatment and environmental exposures. Most complex diseases are the results of a culmination of genetic and environmental factors unique to an individual. Each person's DNA and environmental exposures are unique and assessing both (Figure 2), as well as the interaction of the two, will result in more personalized healthcare. Because nurses are educated on how to assess patients and the environment holistically; nurses are well poised to drive translational research and include information obtained from these assessments to investigating the interaction of environment and epigenomic signatures. Strong communication between bedside nurses and nurse scientists are needed to reconcile the gap between bench studies and what nurses find applicable and useful in the clinical setting.

#### Nursing Policy Implications

Investigation of clinical values that may be directly altered by environmental exposures has implications for nursing policy as well as general public health policies. Person, health, environment, and nursing are the metaparadigm concepts that remain the pillars of the nursing discipline. Perception of these concepts constantly evolves to incorporate new knowledge gained through practice, research, education, and exposure to other disciplines. Since most human disease processes are multifactorial in nature and nurses interact with individuals throughout the illness-wellness continuum, it is imperative nurses become involved with policy development to protect public health

based on research findings that are evident in clinical populations. For example, if the results from this study had indicated that women without GBS colonization had significantly higher serum vitamin D levels additional studies would be needed to validate that the findings were accurate and then to determine if vitamin D supplementation could be protective. After further studies, a significant effort would be required to incite policy change incorporating vitamin D supplementation to prevent GBS colonization into practice. As the state of the science continues to rapidly evolve, it is important to being considering how public health and nursing policies can be addressed and updated to reflect current methodologies and research findings.

#### Nursing Education Implications

Incorporating research models, like the one developed for this study, could have implications for nursing education. Students are usually required to complete basic science courses prior to acceptance and entry into a nursing program (nutrition, chemistry, anatomy, physiology). The base knowledge is required in order to understand how biological phenomena can be utilized, manipulated, and applied to nursing practice. Since the sequencing of the human genome was completed, striking advances in genetics and genomics have occurred and nurses at all levels will be expected to be able to communicate these findings to patients and be able to identify how the gene-environment interactions affect health and illness (Consensus Panel on Genetic/Genomic Nursing Competencies, 2009). Per the recommendations from the Consensus Panel on Genetic/Genomic Nursing Competencies, all programs of nursing should be incorporating education on genetics and genomics for entry level nurses. Essential content that should be taught includes: incorporating genetic and genomic knowledge

into nursing assessment and care, what to do with the information obtained or where to refer patients, understanding how personal opinions of genetic and genomic testing and interventions may affect practice. Translation of genetics into treatment is already occurring and very prevalent in certain areas of nursing, such as maternal-child health and oncology and will become more pronounce in other areas as the state of the science continues to evolves (Kirk, Calzone, Arimori, & Tonkin, 2011). It is imperative that nurses incorporate this knowledge into practice to properly care for patients. Further, "there is a growing abundance of genomic resources already available in a range of formats that cover most teaching environments and learning approaches. For many topic areas, particularly bioscience, there is no need to reinvent the wheel and develop new resources" (Tonkin, Calzone, Jenkins, Lea, & Prows, 2011, p. 336). Content on the epigenetics and appropriate models for analysis should be incorporated into the genomics content due to the increasing number of research studies investigating epigenetic mechanisms contributions to alter health status.

### Limitations

Many of the limitations of this study are a direct result of being a secondary data analysis. Therefore, it is difficult to eliminate the possibility of misclassification bias. The first major limitation of the study is the small sample size. The intent of this exploratory study was to identify significant group differences and determine effect size so that sample size could be determined in future investigations. There have been no previous studies evaluating DNA methylation patterns, serum cytokine levels or vitamin D (25[OH]D) status in pregnant women with and without GBS colonization. Given the high cost of conducting these types of laboratory analyses, conducting a study with a small

sample size to determine feasibility and potential clinical utility is the most economically sound option. Additionally, the intent of using the DAVID database was to identify additional clinically relevant findings that could be utilized as pilot data for a subsequent grant proposal. As outlined in the methods section in the approach for analyzing DNA methylation, both gain and loss of methylation was determined and evaluated. The functional significance varied depending on the direction of change in methylation. We did find significant pathways for methylation gains and losses. However, future studies focusing on genes identified in the methylation loss pathways may prove to have some clinical utility. As discussed previously, the genes involved in the cell morphogenesis and regulation of apoptosis pathways make clinical sense and could potentially be epigenetic factors that contribute to GBS colonization susceptibility.

Another limitation of the study is that only peripheral blood samples were available for analysis and all other information had been previously extracted from the medical record. For example, it would have been more ideal to run the genome-wide DNA methylation analysis on epithelial cells taken from the recto-vaginal swabs at the time of GBS screening because that is when colonization status is determined and that is the reservoir site for neonatal infection. However, we had no access to the swabs or peripheral blood samples at the exact same time point. Since we had maternal DNA samples from the first trimester in pregnancy, any differences associated with GBS colonization could be a clinical indicator for susceptibility. It may also allow for a more targeted screening approach for preventing poor pregnancy outcomes associated with GBS colonization that are currently not prevented or screened. Additionally, a breakdown of the composition of cell types in the peripheral blood samples was not available.

Statistical corrections can be completed to correct for the heterogeneity of cell types found in peripheral blood (Houseman et al., 2012). However, Houseman et al., (2012) states that current statistical correction strategies are "a computationally difficult task that would have extreme vulnerability to model mis-specification." (p. 10). Given the exploratory nature, small sample size, difficulty in establishing model fit, and the fact that the variable cell types that may have an impact on results only make up 2-3% of the cell population, corrections for cell type were not incorporated into the analysis. However, in future studies with a larger sample size consideration will be given to separate cell types prior to analysis of DNA methylation to avoid this type of confounding in the analysis.

#### Conclusions

As the state of the science continues to evolve, it is imperative for nurses to incorporate advances in science into their program of research. Nurses are trained to translate information to people and populations with all levels of understanding. It is a natural fit for nurse scientists to step into a translational role and design studies to assess how the environment interacts with the individual in order to improve health outcomes. Investigation of epigenomic and genomic alterations related to complex disease processes has the potential to identify biologic mechanisms that contribute to the development of disease. Overall, gene-environment interaction models are useful for guiding nursing research investigating DNA methylation patterns because it allows for a holistic approach that clinical assessment data can be incorporated into. Additionally, DNA methylation patterns are readily measurable and offer insight into how environmental interaction can impact health by causing changes in gene expression. A number of standard laboratory protocols and bioinformatics tools can be utilized to complete exploratory studies. Since DNA methylation patterns can now be identified by laboratory techniques, clinicians and research scientists must learn to decipher what the patterns mean and what the implications are for health. As health care progresses to incorporate more patient centered approaches, identification of altered DNA methylation patterns will improve nurses' ability to provide optimal care for patients. With an understanding of the impact of DNA methylation patterns, personalized, individual interventions can be developed to improve health based on research findings. Ultimately, this will improve care at the level of the individual.

APPENDICES

Appendix A

Study Institutional Review Board Approval

UNIVERSITY OF NORTH DAKOTA

INSTITUTIONAL REVIEW BOARD c/o RESEARCH DEVELOPMENT AND COMPLIANCE DIVISION OF RESEARCH TWAMLEY HALL ROOM 106 264 CENTENNIAL DRIVE STOP 7134 GRAND FORKS ND 58202-7134 (701) 777-4279 FAX (701) 777-4708

October 17, 2012

Michelle Lynn Wright Department of Nursing Stop 9025

Dear Ms. Wright:

We are pleased to inform you that your project titled, "Inflammation and DNA Methylation as Group B Streptococcus Colonization Biomarkers" (IRB-201210-092) has been reviewed and approved by the University of North Dakota Institutional Review Board (IRB). <u>The expiration</u> date of this approval is December 31, 2014.

As principal investigator for a study involving human participants, you assume certain responsibilities to the University of North Dakota and the UND IRB. Specifically, any adverse events or departures from the protocol that occur must be reported to the IRB immediately. It is your obligation to inform the IRB in writing if you would like to change aspects of your approved project, prior to implementing such changes.

When your research, including data analysis, is completed, you must submit a Research Project Termination form to the IRB office so your file can be closed. A Termination Form has been enclosed and is also available on the IRB website.

If you have any questions or concerns, please feel free to call me at (701) 777-4279 or e-mail michelle.bowles@research.und.edu.

Sincerely,

Mähelle L Bowler

Michelle L. Bowles, M.P.A., CIP IRB Coordinator

MLB/jle

Enclosures

UND is an equal opportunity/affirmative action institution

#### **REPORT OF ACTION: EXEMPT/EXPEDITED REVIEW** University of North Dakota Institutional Review Board

Date: 10/16/2012		Project Number:	IRB-201210-092
Principal Investigator:	Wright, Michelle		
Department: Nursing			
Project Title: Inflammatio	n and DNA Methylation as (	Group B Streptococcus C	Colonization Biomarkers
The above referenced proje on		nated member for the Ur owing action was taken:	niversity's Institutional Review Board
Project approved. Expe	edited Review Category No must be before:	D	
Copies of the attach must be used in obt	ed consent form with the aining consent for this stu	IRB approval stamp da udy.	ated
Project approved. Exer This approval is valid un periodic review schedule	npt Review Category No tilDecember_31,_20. ed unless so stated in the R	14 as long as emarks Section.	approved procedures are followed. No
Copies of the attach must be used in obt	ed consent form with the aining consent for this stu	IRB approval stamp da udy.	ated
☐ Minor modifications requ approval. This study m	uired. The required correction ay NOT be started UNTIL	ons/additions must be su final IRB approval has	Ibmitted to RDC for review and been received.
	ed. This study may not be		approval has been received.
Disapproved claim of ex Review Form must be fi	emption. This project requir lled out and submitted to the	es Expedited or Full Boa e IRB for review.	ard review. The Human Subjects
Proposed project is not does not require IRB rev	human subjects research as view.	s defined under Federal i	regulations 45 CFR 46 or 21 CFR 50 and
Not Research	🗌 Not Human	Subject	
MUST b	e highlighted and submitte	ed to the IRB within 90	e adviser's signature. All revisions days of the above review date. ducation requirements are met.)
	h	ler in	ACCOLLAN

cc: Denise Korniewicz, PhD, RN, FAAN; Cindy Anderson, PhD, WHNP-BC, FAAN

16/2012 and W Date

Signature of Designated IRB Member UND's Institutional Review Board

If the proposed project (clinical medical) is to be part of a research activity funded by a Federal Agency, a special assurance statement or a completed 310 Form may be required. Contact RDC to obtain the required documents.

(Revised 10/2006)

Appendix B Parent Study UND IRB Approval

Research Project Review and Progress Report	
University of North Dakota Institutional Review Board	

Research Project Review and Progress Report	CHDEVEL
University of North Dakota Institutional Review Board	20 2010 PME
DATE: March 5, 2012 DEPARTMENT: Family and Community Nursing	¥
PROJECT TITLE: Vitamin D Status During Preeclampsia: Mechanisms Underlying Placental Vascular Alterations	LIANCE
PROPOSAL NUMBER: IRB-200809-045	
F MEDICAL COMPONENT, PLEASE GIVE PHYSICIAN'S NAME:	
$\Box$ full board review required, even though original approval was expedited $\Box$ continued approval, "expedited" category $2, 3, 7$	
NEXT REVIEW REQUIRED BEFORE: <u>March 22, 2013</u>	
$\Box$ CONTINUED APPROVAL, BASED ON FULL BOARD REVIEW	
APPROVAL TERMINATED COMMENTS OF REVIEWER:	
	_
c: Chair, Family and Community Approval Date: 2-13-12-	-
Nursina	]
I. Is project complete? Yes □ No ⊠	
2. Is project ongoing? Yes ⊠ No □ If No, explain below and indicate if continued approval and continuing review is desired.	
3. How many subjects have been enrolled in the research project?	
• •	
since the date of last approval, and	
$\frac{1}{l_{e}5}$ since the date of last approval, and since the initial approval	
$l_{e}5$ since the initial approval	
$l_{e}5$ since the initial approval	
$l_{\rho}$ since the initial approval 4. Is the research permanently closed to the enrollment of new subjects? Yes $\boxtimes$ No	
<ul> <li><u>(e5</u> since the initial approval</li> <li>4. Is the research permanently closed to the enrollment of new subjects? Yes X No □</li> <li>Have all subjects completed all research-related interventions? Yes X No □</li> <li>Does the research remain active only for long-term follow-up of subjects? Yes X No □</li> <li>5. Is data analysis complete? Yes □ No X</li> <li>*** If the research is permanently closed to the enrollment of new subjects, all subjects have completed all research-related nterventions, the research does not need to remain active for long-term follow-up of subjects, and all data analysis is complete,</li> </ul>	

- 6. Has any additional grant money been awarded for this project in the past year? Yes No K If yes, submit a copy of the grant along with this completed form.
- Describe any adverse events and/or unanticipated problems involving risks to subjects or others that have occurred since the last approval. If you did not report the adverse event or unanticipated problem previously, a separate Unanticipated Problem/Adverse Event Form must be submitted to RD&C with this form.

No adverse events or unanterpated problems

Have any additional risks with this research been identified? Yes No K
 Describe all benefits experienced by participants, and include a current risk/benefit assessment based on study results.

Participates expressed extrusion about their contributions to generation of new Knowledge to insprove pregnancy surcordes

- 9. Have there been any changes or deviations from the approved protocol since the most recent approval? Yes No K If Yes, elaborate below, and submit a separate Protocol Change Form to the RD&C indicating proposed protocol changes.
  - a. Have any of these changes been implemented already? Yes No If yes, please describe fully.
  - b. Are any protocol changes being planned for later implementation? Yes No A lf yes, please describe fully. A separate Protocol Change Form must be submitted to RD&C for approval before the proposed protocol changes can be implemented.
- 10. Have any subjects withdrawn from the research? Yes 🖄 No 🗌 If yes, state how many have withdrawn and describe the circumstances.

I have wethdrown due to note away from study site and pregnancy termination. None withdrown since last continuing renew

Research Project Review and Progress Report

10/10/07 2 11. Have there been any complaints about the research since the last IRB review? Yes 🗌 No 📈 If yes, please report and summarize the complaints and your response/action.

12. Summarize any multi-site trial reports relevant to your research.

13. Summarize any recent literature, findings, or other information relevant to your research, especially

information about risks associated with the research. No works reported, fourcasing evidence regarding health works in vitament D definition.

- 14. Have all/PI's involved with the research completed the IRB Educational Requirements? Yes 🗹 No 🗌 (Educational requirements must be completed before the IRB can grant continued approval for the research project.)
- 15. On a separate piece of paper, provide a thorough protocol summary (approximately 300 words) giving a concise summary of the protocol's progress to date and the reasons for continuing the study or reasons for asking the IRB to terminate approval. The summary should include, for instance, an explanation of any complaints about the research, relevant multi-site trial reports, participant benefits, or a current risk-benefit assessment based on study results. Sufficient information is required in the summary so that the IRB can determine whether the proposed research continues to fulfill the criteria for approval.

attached

16. A copy of the current informed consent document(s) (with the IRB Approval stamp), as well as a clean copy of the consent document(s) (with no IRB Approval stamp) must be submitted with this report.

Current copy included. Clean copy included

17. Have there been any changes in the conflict of interest statement or situation for the Principal Investigators, research staff involved in the study, or each individual's respective family members in the last 12 months? Yes D No Z If yes, please describe fully on a separate sheet of paper.

Signature of Principal Investigator Cindy MandeusenDate 3/19/12
Signature of Principal Investigator <u>Circly Mardusmate</u> <u>3/19/12</u> Current email address: <u>Cindy</u> , anderson @ email. und. edu
Current Address: Stop 902.5

This completed form should be returned to the IRB, University of North Dakota, 264 Centennial Drive Stop 7134, Grand Forks, ND 58202-7134.

Research Project Review and Progress Report

10/10/07 3

Appendix C Parent Study Altru IRB Approval

		Institutional	Review Boar	d (IRB)	5/24
HEALT	H SYSTEM	Contin	uing Rev	iew	5 [-
					Revised 5/10/11
Date: E	5/16/12	Anderson	IRB #	ST-54	
Principal Inves	ligator: Cindy	Anderson			
Department:	Nursina		Phone #	777-	4354
Research Coord	linator:	х.	Phone #		
Project Title:	Vitamin D Status dur Alterations.	ing Preeclampsia: Mech	anisms Underlying	Placental V	ascular
	TOR COMPLETE: Expedited Review?			X	Yes No
	-	918108			
<ol> <li>Date of</li> <li>Date of</li> </ol>	Last Approval: (if applica	able) 4/11			
		h done, enrollment close &			
		If no, complete questions 5			
5. Is the r	esearch permanently close	ed to the enrollment of new	subjects?	······ <u>×</u>	Yes No Yes No
6. Have a	Il participants completed a	all research-related interven	tions?	×	_ 105 100
		gress of the project (i.e., nu to go on, etc.) Classed M Wedded d to gache m the research?			
expec	ted to exte	nd to gure	2013.		V., No
8. Have a	ny subjects withdrawn no				
9. Have	e with draw called the complete	Adrawn and describe the circ M = 8 deal M = 2 Mes ca M = 4 / 1 La (concerns about the research	int competents int to call since the last IRB rev	char char no - ce	ged prace
If yes,	please report the complair	nts/concerns and your respo	onse/action.		
10		olving risks not detailed in	your original applicativ	n <sup>2</sup>	Yes No
10. Any u 11. Any s	nanticipated problems invo erious adverse events (SAF	E) at your site?	your original application		Yes No
11. AUV S	AE, safety update, action l			-	

If yes to any of the above (7-13), please provide a detailed summary below. Need to write a more detailed summary about the adverse events, problems, etc. (i.e., the types of adverse events, numbers, trends).

(Example: There were 6 adverse events during the past year, four of which occurred in patients at other sites and 2 occurred at this site. Three of the events were constipation thought to be related to the study drug; two were shortness

Page 1 of 2

nn	eath thought not to be related to the study drug; and one was a petechial rash thought to be possibly related.)
КB	Continuing Review
Also, r	make certain adverse events/mortality reports have been submitted and are on file in the IRB office.
4.	Any protocol changes, consent form changes or amendments?
5.	Number of subjects enrolled since last review:
16.	Total number of subjects enrolled since project initiated: <u>67</u>
17.	Have all PIs involved with the research completed the IRB Education Requirements?
18.	(Education requirements must be completed before the IRB can grant continued approval for the research project) A copy of the current informed consent document (with, if possible, the IRB Approval dated stamp) must be submitted with this report.
	, ,
Signat	1 Charles 5/16/12 thure of Principal Investigator Date
****	Date
****	Date
****	Date
****	Date Date USE ONLY:
****	Date         ************************************
IRB	Date         USE ONLY:         ✓         Continue approval based on Expedited Review         θ         Continue approval based on Full Board Review         θ         Suspend approval, pending investigation         MMMM         MMMM
IRB	Date         EUSE ONLY:              Continue approval based on Expedited Review            Continue approval based on Full Board Review            Suspend approval, pending investigation

Page 2 of 2

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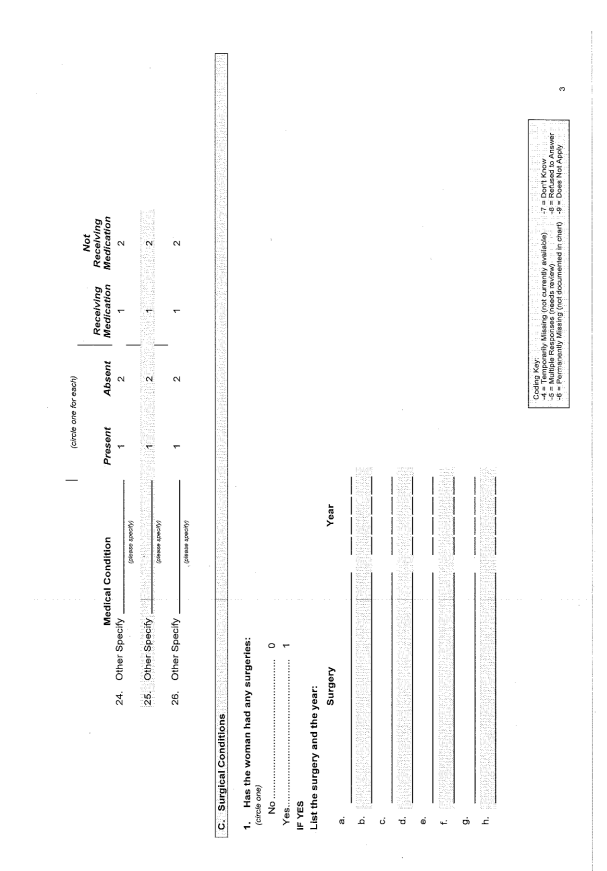
Appendix D

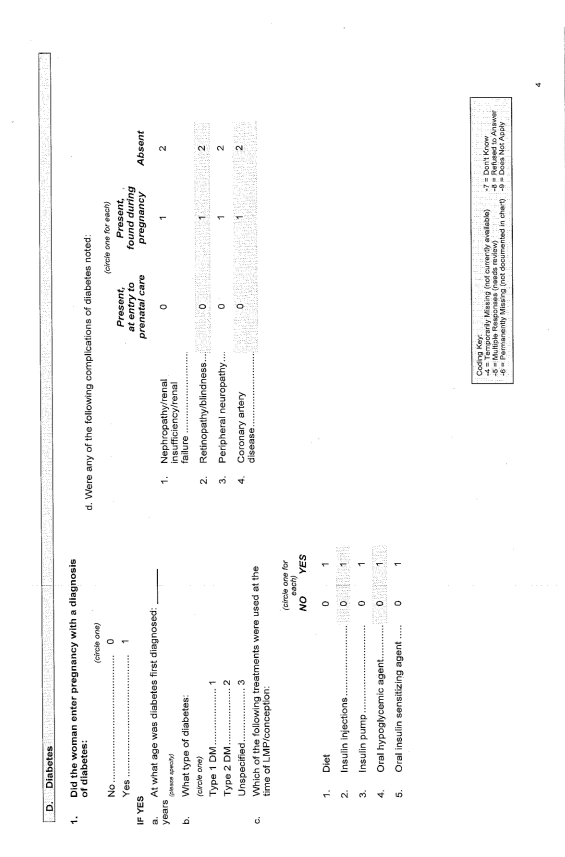
Medical Record Abstraction Form

Date of Death:  $-\frac{J}{mm \frac{2dJ}{M}}$ If LB & Child NOT Living Coding Key: = Temporarily Missing (not currently available) = Temporarily Missing (not currently available) = Temporarily Missing (not documented in chart) = Permanently Missing (not documented in chart) = Permanently Missing (not documented in chart) Reason for death: Maternal Age at Time of Delivery: Code: Child NOT Living (Reason for Death) 01=Carciovascular 01=Carciovascular 03=Respiratory 04-Prematurity 06-Unknown 93=Cuther ± |\_\_\_ 2) [\_\_\_\_] 3) [\_\_\_\_] (5 Code Fetal/Infant **ID Number:** Description CO-Nome noted CO-Nome noted C2. Small for Ceretational Age C2. Small for Ceretational Age C4. Terge for Ceretational Age C4. Terge for Ceretational Age C4. Terge for Caretational Age C4. Terge for Caretational Age C4. Terge for Caretational Caretational C4. Cheretational Fetal/Infant Complications Birth Weight Ę ลิ ଳ 3)[\_\_\_\_ 2) Code Complexite mode Comple Maternal Maternal Complications Complete this medical chart abstraction form for each participant enrolled in the study. Description Date of First Prenatal Visit: ç ลิ ଳ l Ö ŝ Type of Delivery grams 01=vaginal 02=op vaginal (forceps or 03=C-section Type of Delivery ົຈ 01≕male 02≃female Gender ---| ---Enter time in weeks/days if documented. 01 fit number of weeks is not documented. of the set of t GA (Gestational Age) code A. Detailed Past Obstetrical History ₹ GA: |\_\_\_\_\_/ / |\_\_\_| wks days ËDC Gender: | ∺ Pregnancy Outcome Time: **Gravida Status** LB=Live birth SB=Stillbirth SA=Spontaneous Abortion SA=Spontaneous Abortion TA=Therapeutic Abortion TP=Tectopic Pregnancy MP=Molar Pregnancy ġ. Reviewed by; Outcome: Instructions GA code: Study 608 LMP: \_\_\_\_ Ċ. Outcome

		(circle one for each)	for each)	•	:
	Medical Condition	Present	Absent	Receiving Medication	Not Receiving Medication
÷.	Asthma	-	5	-	2
2	Seizure disorder	V	N	Ł	7
ကံ	Chronic hypertension	~	Ю	~	ы
4	Diabetes mellitus (type 1 & 2)	2	N	÷	8
ъ.	Hyperthyroidism	-	7	4	N
Ó	Hypothyroidism	Ł	R	٢	N
7	Valvular heart disease	1	2	-	0
œ	Other structural heart disease	Ł	0	Ţ	7
ດ່	Coronary artery disease/				
		£	2	-	0
ę	. Nephropathy/Nephrotic syndrome/				
	Glomerulonephritis	~	N		8
-	. Renal insufficiency/ renal failure	-	ы	۰	ъ
2	Sickle cell anemia	<b>Χ</b>	7		8
7		~	N	~	0
Ţ	. Lupus erythematosus	Ļ	N	-	Ŋ
÷	<ul> <li>Antiphospholipid antibody syndrome</li> </ul>	~	0	~	7
9	. Rheumatoid arthritis	4	N	Ţ	8
17	. Ulcerative colitis/Crohn's disease	~	2	~	ю
19	<ul> <li>Malignancy Specify (please specify)</li> </ul>		N		8
20.	. Hepatitis B	٣	5	~	7
2	Hepatitis C		3		2
22.	Psychiatric Disorder Specify	1 1	2	~	7
50	Other Specify		2		2

N





Cooling Keyr. 4.4 Temporanity Missing (not currently available) 5.5 Milliple Responses (needs review) 9.5 E Permanently Missing (not documented in chart) 9.5 E Does Not Apply Vaccines (600 series) 610=Influenza 630=Rtepatitis B 630=Rubella 640=Varicella-zoster immune globulin (VZIG) 699=Other vaccine Vitamins (500 series) Postpartum Th<u>vroid Agents</u> 251=Antithyroids (overactive) 252=Thyroid Replacement (under active) 399=Other Medication Trimester 01=Taken 02=Not taken 1<sup>st</sup> 2<sup>nd</sup> 3<sup>rd</sup> 170-Antiemetics
 180-Antipsychotics
 190-Antivirals
 200-Birth control pills
 210-Chemotherapeutics
 220-Diuretics
 230-eff agents
 240-Progesterone
 260-Rhogam
 270-Steroids E. Prescription Medication, Vitamins and Vaccines other Antihvpertensives 1661 = Antihvpertensives 1621 Labetolol 1638 = Labetolol 1638 = Ca-Channel Blockers 1645 = Beta-Blockers 1655 = Ace-Inhibitor 1655 = Ace-Inhibitor 1658 = Ace-Inhibitor 1658 = Ace-Inhibitor Medication, Vitamin, Vaccine Code\* 120=Anticoagulants130=Antidepressants140=Anticonvulsants150=Antihistamines 110=Antibiotics Name Analgesics 101=Narcotic 102=NSAID 103=Aspirin 104=Acetaminophen Medications (100-300 series) <del>1</del>0. Ň റ് ÷ 4. œ N പ് ы. ۍ

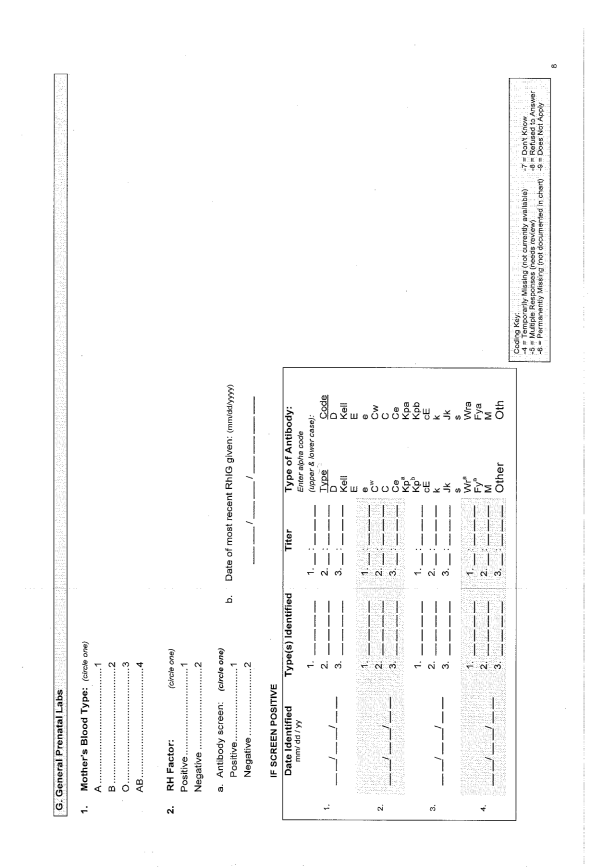
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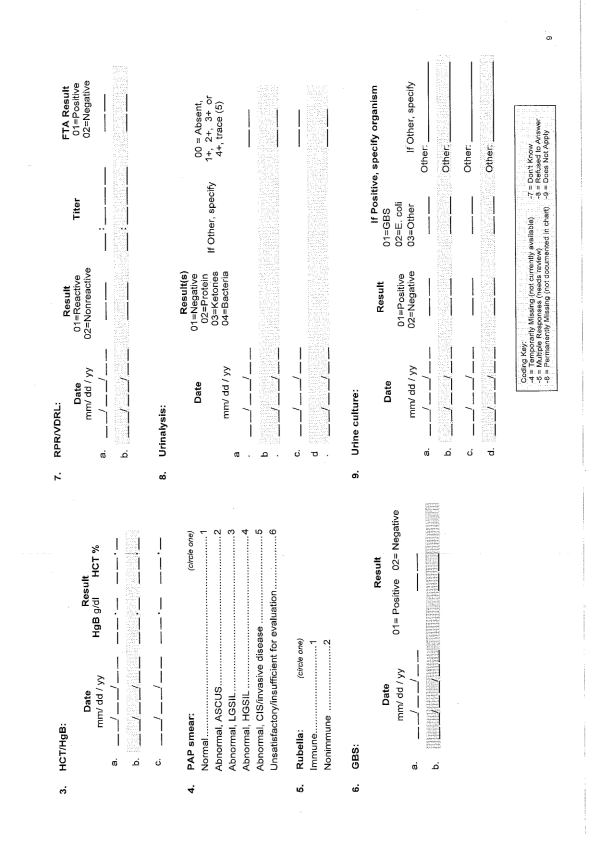
G Coding Key. 4 Temperatry Missing (not currently available) 5 Multiple Responses (needs review) 6 Retused to Answer 9 Permanently Missing (not documented in chart) 9 Poes Not Apply Other Condition Complete the Prenatal Care Flow sheet on all women enrolled in the study using medical records. Complete one row for each prenatal visit. Specify 01=Negative 02=Trace 03= > +1 Urine Dipstick Glucosuria 01=Negative 02=Trace 03=+1 04=+2 0 05=+3 06=+4 0 Urine Dipstick Proteinuria 00=normal 01=decreased 02=absent Fetal Movement Fetal Heart Rate 01=present 02=absent Fundal Height E 1. Sys\_\_\_\_\_\_ 2. Dia\_\_\_\_\_\_ Highest Blood Pressure mm Hg 2. Dia 1. Sys\_\_\_\_\_ 2. Dia 1. Sys\_\_\_\_\_ 2. Dia\_\_\_\_ 1. Sys\_\_\_\_\_ 1. Sys\_\_\_\_\_ 2. Dia\_\_\_\_ 1. Sys\_\_\_\_\_ 2. Dia\_\_\_\_\_ 1. Sys\_\_\_\_\_ 2. Dia\_\_\_\_ 1. Sys \_\_\_\_ 1. Sys\_\_\_\_\_2. Dia\_\_\_\_\_ Weight sq F. Prenatal Care Visits Ì | | | l mm/ dd / yy Date | | Instructions İ h Ì Ì Ì 1 1 Visit ÷ ۲. ő 6 ni ë. 4 ю 6

Other Condition	Specify									
Urine Dipstick Glucosuria	01=Negative 02=Trace 03= > +1									
Urine Dipstick Proteinuria	01=Negative 02=Trace 03=+1 04=+2 05=+3 06=+4									
Fetal	Movement 00=normal 01=decreased 02≖absent									
Fetal Heart										
Fundal	Height	5								
Highest	Blood Pressure	1. Sys 2. Dia	1. Sys	1. Sys 2. Dia	1. Sys 2. Dia					
Weight	<u>4</u>	8								
Date	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				//			//		//
Visit		10.	1.	4	33	14,	15.	16, 1	17.	18

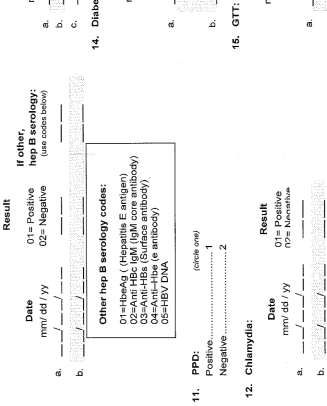
Coding Kay: 4 = Temporarity Missing (not currently available) 5 = Multiple Responses (needs review) 6 = Permanently Missing (not documented in chart) 9 = Does Not Aptily

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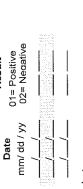


10. Hepatitis B Surface Antigen:



13. Gonorrhea:

Result



ю. Ю ΰ 14. Diabetes screen:

sult	Random		🗆 mg/dl		
Blood Sugar Result	1 Hour	······	Ip/gm		
	Fasting		🗆 mg/dl		
Date	mm/ dd / yy			. 1 1	

□ mg/dl □ mmol/L

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□ mg/dL

🗖 mg/dl

D mmol/L 3 hr □ mmol/L Blood Sugar Result 2 hr D mg/dl 1 hr ID/gm [] Fasting mm/ dd / yy Date

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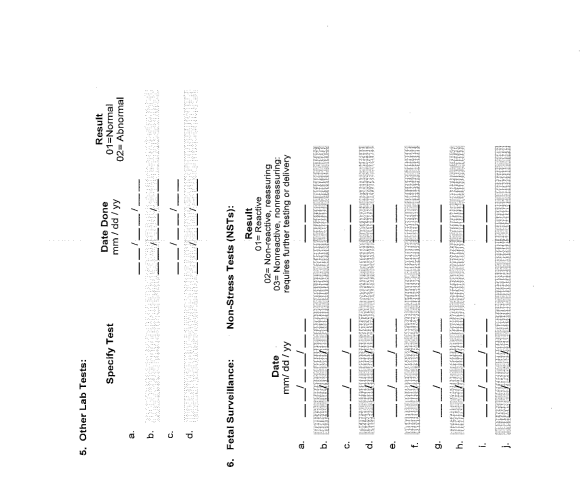
10

H. Specialized Testing					
Instructions					
Complete the Specialized Prenatal Labs on all women enrolled prospectively into the study who have results of specialized testing in their charts. All questions refer to the mother. If the number of tests done of a type exceeds the number of rows provided, enter the results from the first prenatal lab tests on the first row. Use the remaining rows to enter the most recent results chronologically.	y into the study w ber of rows provide	tho have results ed, enter the res	of specialized ults from the	testing in their first prenatal lab	charts. All questions tests on the first row.
1. Specialized testing: (circle one)	4. Pre-Eclampsia Labs:		-		ò
No	Test	Date Done mm/dd/yy	Result	Units	
2. Hemoglobin (HgB) A1C:		1 1		⊇	
Date Result mm/ dd / yy %		1 1		IU	
<b>b</b> .	HQ	1 1		D	
		1 1		וח	
3. Twenty-four hour urine protein:				D	1
Date Result mm/ dd / yy mg/24 hr	AST	1 1		D	
	(SGOT)	. / /	-	Ŋ	
		1 1		⊇	
		1 1		Ð	
	ALT	/ /		C	
	(SGPT)	1 1		D	
		/ /			

÷

		5000 E				Urine	Prot/ Creat Ratio				24- hour urine	protein	
1 1	1 1		1 1	Date	mm/dd/yy			-	1 1			· ·	1 1
				Ratio	Result					Result			
K/uL	Kul	Kut	Kut			urine total protein mg/dL urine creatinine, random mg/dL	urine total proteinmg/dL urine creatinine, randommg/dL	urine total proteinmg/dL urine creatinine, randommg/dL	urine total proteinmg/dL urine creatinine, randommg/dL	mg/24 hr	mg/24 hr	mg/24 hr	mg/24 hr

mg/dL	mg/dL	mg/dL	mg/dL	mg/dL	mg/dL	mg/dL	mg/dL	mg/dL	mg/dL	mg/dL	mg/dL	g/dL	g/dL	g/dL	g/dL
												-			
1 1	1 1	11	1 1	1 1	1 1	1 1	1 1	I $I$	I I	1 1	1 1	1 1	1 1		1 1
		Uric Acid				Z				Creatione				Паподолан	



	And and a second s							
			Measurements		Kesult			
	Date		CRL=Crown-rump length BPD=Biparietal diameter HC=Head circumference AC=Abdominal circumference	Code (see codes on page 17)	Specify	AFI (Amniotic Fluid Index) cm	BioPhysical Profile	
			Consistent with:				Fetal Tone	12
		1. CRL	cm wks gestation	     			Gross body movements	12
-		2. BPU 3. HC	where the set of the set o	5 <del>4</del>     		Check If	Fetal breathing movements	2
<u>.</u>		4. AC 5. FL		2.		ent is	Amniotic fluid volume	12
+	-		Consistent with:			-	Fetal Tone	12
l	///	1. CRL	cm wks destation	5			Gross body movements	12
		2. BPD		3.		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Fetal breathing movements	2
N		ν. 4 Ο Ο Α.	which are the set of the se	2 <sup></sup>		measurement is	L vieckin measurement is Amniotic fluid volume	1
		5. FL	cm wks gestation				-	2
			Consistent with:	1. 			Fetal Tone	/2
		1. CRL	cm cm wks gestation	2.			Gross body movements	12
		2. BPD	cm wks gestation	э. •			Fetal breathing movements	/2
		4. AC		22		measurement is normal.	measurement is Amniotic fluid volume normal.	12
		5. FL	cm www.gestation			:	-	
			Consistent with:	1.			Fetal Tone	/2
		1, CRL 2. BPD	cm wks gestation	3.			Gross body movements	2
4.		3. HC	cm wks gestation wks gestation	4		Check if measurement is	Fetal breathing movements	12
		4. AC 5. FL	cm wks gestation			normal.	Amniotic fluid volume	5

I. Prenatal Ultrasound

L

	AFI (Amniotic Fluid Index) cm	Eetal Tone     Fetal Tone     Gross body movements     Gross body movements     Gross body movements     Fetal breathing movements     Fetal breathing movements     Amniotic fluid volume     Commal.	Fetal Tone2 Gross body movements2 Fetal breathing movements2 measurement is Amniotic fluid volume2 normal.	Fetal Tone     /2       Gross body movements     /2       Check If     Fetal breathing movements     /2       measurement is     Anniotic fluid volume     /2       normal.     Fetal Tone     /2	/ movements hing movements
Result	Code (see codes on page 17) Specify	9. 4. 9. 7. 9. 4. 9. 1	5	, φ,	
Measurements	CRL=Crown-rump length BPD=Bipartetal diameter HC=Head circumference AC=Abdominal circumference FL=Femur length	Consistent with: cm wks gestation cm wks gestation cm wks gestation cm wks gestation cm wks gestation	Consistent with: Consistent w	Consistent Consistent	Consistent wut: Cm Cm wks gestation Cm wks gestation Cm wks gestation Wks gestation
	Date	4. AC	<b>5.</b> FL	<b>6</b> <b>6</b> <b>7</b> <b>1</b> <b>1</b> <b>1</b> <b>1</b> <b>1</b> <b>1</b> <b>1</b> <b>1</b> <b>1</b> <b>1</b>	7. 1. CRL 3. HC 7. 5 BPD 7. 6 FL 7. 7 CRL 7. 7 C

*Results/Interpretation of Ultrasound		
100=No structural abnormalities 101=Dates consistent with EDC	Structural anomalies:	<u>Structural anomalies (Continued)</u> Abnormal placenta
OB complications: 110=IUGR/SGA (Specify growth %-tile)	<u>Markers</u> 301=Choroid Plexus Cyst (CPC)	321=Previa 322=Abruption / sub-chorionic hemorrhage
120=Macrosomia (Specify growth %-tile)	302=Pyelectasis (4 -10 mm) 303=Fchogenic Intra-Cardiac Focus (ECF)	329=Other (specify)
140=Polyhydramnios (AFI ≥ 25)	304=Ventriculomegaly (≥10 mm) 305=Bound humarachoic	Abnormal Cord insertion 331=Marcinal
ן זטט≓כפרעוכפו snortening (> בס ווווו פו >∠ס שפראכ) לל60=Fetal arrhythmia	306=Shortened femur or humerus	332=Velamentous
170=Fetal demise	307= Increased nuchal translucency (first trimester)	339=Other (specify)
Multiple gestations:		Abnormal Umbilical cord
200=Twins - di/di	Congenital Anomaly	341=ZVC
210=Twins - mono/di	310=Congenital Anomaly	04Z=NUCTAI
220=Twins - mono/mono	Under the specify column, note the code	040-Varix 040-Other/energia/
230=Triplets or higher order gestation	from the congenital anomaly code list	oda-Outer (specify)
240=Concordant growth (multiple gestation) -All ≥10%	on the following page.	350 = Amniotic Band
250=Concordant growth multiple gestation) -All SGA (<10%)		
260=Discordant growth (multiple gestation) – At least one SGA (STU20) 270=Dercent discordant(Shecify %)		<u>Other</u> 499=Other (specify)
(calculated by Big twin-Little Twin Weight/Big twin)		

Exr Dosuit Code 310: Specify Condenital Anomaly Code	111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111 - 111	
100=Cvstic Hvaroma	Congenital Heart Defects	Genitourinary defects
	401=Atrioseptal defect (ASD)	601=Hydronephrosis/ureteropelvic
Central Nervous System Defects	402=Ventricular Septal Defect (VSD)	junction obstruction
201=Open neural tube defect (Meningomyelocele, Spina	403=Atrioventricular Canal Defect	602=Autosomal Recessive Polycystic
bifida)	(Endocardial Cushion Defect)	Kidney Disease
202=Ánencephaly	404=Transposition of the Great Vessels	603=Mutlicystic/Dysplastic Kidney
203=Hydranencephaly	405=Tetraloov of Fallot	604=Posterior Urethral valves (PUV)
204=Hydrocephalus	499=Other Congenital Heart Defect (Specify)	605=Renal Agenesis
205=Holoprosencephaly		699=Other GU defect (Specify)
299=Other (Snecify)	Gastrointestinal defects	
	501=Cleft Palate	Skeletal defects
Thorax Defects	502=Gastroschisis	701=Skeletal dysplasia
301=Congenital Diaphragmatic Hernia	503=Omphalocele	702=Club feet
302=Cystic Adenomatoid Malformation (CAM)	504=Duodenal Atresia	799=Other limb abnormalities (Specify)
303=Pulmonary Sequestration	599=Other Gastro-Intestinal Defect (Specify)	
399=Other (Specify)		Otter 600-Other anomaly (Snacify)

	End date:	End time						
-	If yes: Start date:	Start time:						
On MgSO4? (circle one)		e e	Kes	о с	Kes	ğ	Ves	or
Highest Blood Pressures	1. Sys	7. Da	1. Sys	2. Ua	1. Sys		1. Sys	
Date		//				—		— — / /- —

Appendix E

DNA and Primer Sequences for Validation

# CpG Site DNA Sequence Data for Primer Design

RHPN1>hg19\_dna range=chr8:144457427-144457927 5'pad=250 3'pad=250 strand=+ repeatMasking=none

HLA-DRB6>hg19\_dna range=chr6:32522622-32523122 5'pad=250 3'pad=250 strand=+ repeatMasking=none

ANXA2>hg19\_dna range=chr15:60643907-60644407 5'pad=250 3'pad=250 strand=+ repeatMasking=none

GGCCACATTCACTTACCCAGGTTCAGGAAAGCATTTTCCAGGTCTCCTTTAAC CTCTTTCCTGATGCTTTCCAACATGTCATAAGGGCTGTAACTCTTGTACCTATC AAATACTGAGGAAAAACAACAAAGAGTTATCAGATCCGAGCCACTAGTCAA AGCTGTCAACGATCACCCACCTAGTTTTATGCACCATAATTTTTTTAAAAATT GAGGATGATCACAGCATCCTAGGAGCTTAGAGGTTACCACGGTGACCAGAGC CAACATTGGCCAAGTTTGTCGTGGAACAGCCATACCACCTGTCCTGAATGGC ACTGCCCAGGCCACATATTTGGACCATCTCTATCTCCCCTGAGTGGAACCAT TCCATCCGAAAACCATAGGAAACAGTACAGAGCATGCACCAAAGTCCACTAC TTCAACAAATAATGGCAAGACCAAATGATCATCAAACAAGAAGGAGCTGCA GAATAAAGCACCAAATGCAGAAACTATTTG

MRI1>hg19\_dna range=chr19:13874611-13875611 5'pad=500 3'pad=500 strand=+ repeatMasking=none

CCTCAGCCTCCCGAGCAGCTGGGACTACGGGTGTGCGCCACCACGCCCAGCT AATTTTTTGTATTTTAGTAGAGAGACTGGGTTTCACCATGTTGGCCAGGCTGGT CTCGAACTCCTGACCTCAAGTGATCTGCCCGACTCGGCCTCCCAAAATGCTGG CGAGCAGAAAATGGCCCAGAAACAGCCTTGCATCCATCAAGGGGCACACGA CCCCCCACTACCTCCCCCTCAACCTTGGAAGATCATTTAACAAATTCTTTGGT TTGAACACTTGATGTTACCTTGCCACTGGGGATACATCCCTAACTCTAGACAG CAGGTTGTTAAACACGGGGCCTGGTATCCACTAGGCGTCCCATAAATGCTGC CACTTTTGTGGTTCCGAGGAGGCGGCTCACTCCGTTCGGGCTTGGCAGGAGTC GTGGAGTGGGTTCGGCCACGTGGAATCCGCGTCCTGGGAACCCGTGGAATCC GCGTCCTGGGAACCCGTGGAATCCGCCTCCTGGGAACCCGTGGAATCCGCCT CCTGGGAACCCGTGGAATCCGCCTCCTGGGAACCCGTGGAATCCGCCTCCTG GGAACCCGTGGAATCCGCCTCCTGGGAACCCGTGGAATCGGGTTGGATGCGC ATGTGCGTGTCTCTTTTTCCGGGGGGGGGGGCCCCGCCCACGGCCCCGCCCCGCTC CCAAGTGCGCGCGGACCCCTAGCTCCCTCTGAGTTGCGCTGGGCTTGGCTGCT GCACCATGACCCTGGAGGCGATCCGCTACTCGCGGGGGCTCCCTGCAGATCCT AGACCAGCTGCTGCCCCAAGCAGAGCCGCTACGAGGCGGTGGGCTCGGTG GGCGGGGGCG

GAP43>hg19\_dna range=chr3:115376099-115376599 5'pad=250 3'pad=250 strand=+ repeatMasking=none

AGTGTAGGAGAGGTGAGTTGCTTAGGTCTAAGGAGAAAGACTGCTTAGGTGT GTGTTCACCCCCAGGACGAAGAAAGGAACACTGGGTGAGATTTTGTTCAACT ACCCATAGTTACCACCAGATGGTGAAACTGATCCCGGGCCTCTTGGGTATTG ATCAGTTTATGGGGAGATGGGGGAGAAGACTATCTTTCACTTGTTAATTCATTA ATTTCTTTCGCAAATATTTTTTCAGTACCTGCTAAGTCCCACGGACTATGCTA GGAGCTGCTGTTAAAATGACAAACCAGATAAGGTCACTGCCCTTAATCAACT TACAGTTGGGTGAGAAGCTATCAGGTACAAGTATGGCCCTAGAACAAATTAG TCTTTTCTAGTTAATAATCTTATGTGATGAGATTTGGCCTTGCTCCTTTGGTGA CTTGCCTCAAGGAGCCCCAGGCAAAACCAATGTAACATATATTAATAATATA TGAAATAATATTTTTGTAGACACAATG

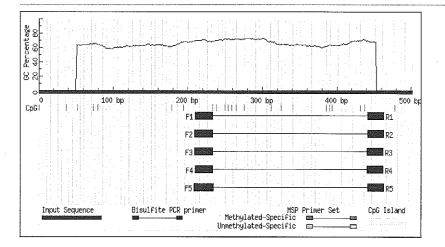
CUL3>hg19\_dna range=chr2:225441582-225442082 5'pad=250 3'pad=250 strand=+ repeatMasking=none

ATCATATATGAACTTCTGTTTTTGAAGCCACCCCTCAAGAGCCAACAGGATTC TTTAAGTATCCCAGTGGTACTAAACCCATATCCTTTGAGAATGCTTTCTAGA ACGATTCACAAACTGGCTCTGCAGGCTTTTCAAAACTTAAGTTCTAGAAGTTG TACAATAAAATGACAGGATCGCTAAAATAAGTGTATGGCATTCGATGTAACT GCTTGCAAAAACAACATCCAATTTTAATATTGGCCTAATCGTGGCTAAATATT GGTATAATAATAGTTAACTATCTGCTAAGTTCTATTTTAAAGCTTTATTTTATT TATCACACAACAACCTGTAAGGTGGGTCTCATTAGCAGTCTCATTTTACACA GGGGAAAACTGAGGCTACAAGAAGTAACTTGTTAAAGGTTATGCAGCTAGAG GCCGGGCGCGGTGGCTCACGCCTGTAATCCCAGCACTGTAGGAGGATGAGGC AGACGGATCACGAGGTCAGGAGATCG

#### MethPrimer Results - MethPrimer - Li Lab, UCSF

### MethPrimer result

Please cite MethPrimer: Li LC and Dahiya R. <u>MethPrimer: designing primers for methylation PCRs</u>. Bioinformatics. 2002 Nov;18(11):1427-31. PMID: <u>12424112</u>



Sequence Name: Sequence Length: 501

CpG island prediction results (Criteria used: Island size > 100, GC Percent > 50.0, Obs/Exp > 0.6) No CpG islands were found in your sequence Primer picking results for bisulfite sequencing (or restriction) PCR

	Primer	Start Size	e Tm	GC∛	'C's	Sequence
1	Left primer	210 24	58.03	62.50	7	GGATGTATTTTTTTTAGTGGTTGG
	Right primer	463 22	58.45	72.73	6	CCTCACCCAAATAAACCCTACT
	Product size: 254	, Tm: 69.8,	CpGs in	product	: 15	

2 Left primer 209 25 58.88 60.00 7 AGGATGTATTTTTTTAGTGGTTGG Right primer 463 22 58.45 72.73 6 CCTCACCCAAATAAACCCTACT Product size: 255, Tm: 69.9, CpGs in product: 15

- 3 Left primer 209 25 58.88 60.00 7 AGGATGTATTTTTTTAGTGGTTGG Right primer 462 22 56.82 72.73 6 CTCACCCAAATAAACCCTACTC Product size: 254, Tm: 69.8, CpGs in product: 15
- 4
   Left primer
   210
   24
   58.03
   62.50
   7
   GGATGTATTTTTTTTAGTGGTTGG

   Right primer
   462
   22
   56.82
   72.73
   6
   CTCACCCAAATAAACCCTACTC

   Product size:
   253, Tm: 69.7, CpGs in product:
   15
- 5 Left primer 208 26 58.92 61.54 8 TAGGATGTATTTTTTTAGTGGTTGG Right primer 463 22 58.45 72.73 6 CCTCACCCAAATAAACCCTACT Product size: 256, Tm: 70.1, CpGs in product: 15

61 CAGCTTACCCCACCGACCACGTCCTTCTGCATTGACTGCCTCCTGTCCTGCCCAG

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

12/8/13

12/8/13

301 GTGACTCCCTGACGCAGATCCAGTGCGGCCAGCTGCAGAGCCCCAGGGCCCAGATTCACC ||||:|::::||++:||||::|||++|::||:||:|++:||||::::|||||:::: 301 GTGATTTTTTGACGTAGATTTAGTGCGGTTAGTTGAGAGTCGTAGGGTTTAGATTTATT

481 CAGGCAGATGTCTGCCCCATG :||:||:||:|:::::|| 481 TAGGTAGATGTTTGTTTTATG

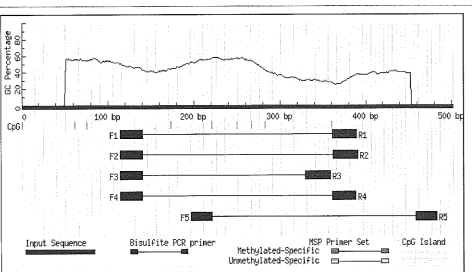
MethPrimer v1.1 beta

Li Lab, Department of Urology, UCSF

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

#### MethPrimer Results - MethPrimer - Li Lab, UCSF

## **MethPrimer result**



Please cite MethPrimer: Li LC and Dahiya R. <u>MethPrimer: designing primers for methylation PCRs</u>. Bioinformatics. 2002 Nov;18(11):1427-31.

PMID: <u>12424112</u>

12/8/13

Sequence Name: Sequence Length: 501

CpG island prediction results (Criteria used: Island size > 100, GC Percent > 50.0, Obs/Exp > 0.6) No CpG islands were found in your sequence Primer picking results for bisulfite sequencing (or restriction) PCR

1	Primer Left primer Right primer Product size:		114 389	26 28	56.17	42.31 35.71	4 4	Sequence TATTTTAGGATGGATTAGGAGAAAAA CAAAAATTTATAAACACTTCAACAATAC
2	Left primer Right primer Product size:	277,	390	29	57.95 57.60 CpGs in	34.48	4	TATTTTAGGATGGATTAGGAGAAAAA TCAAAAATTTATAAACACTTCAACAATAC
3	Left primer Right primer Product size:		359	30	57.95 57.71 CpGs in	26.67	4	TATTTTAGGATGGATTAGGAGAAAAA AACAATTTCAAACTATAAAAAAAACAAATAA
4	Left primer Right primer Product size:	275,	388	27	57.95 53.96 CpGs in	33.33	4	TATTTTAGGATGGATTAGGAGAAAAA AAAAATTTATAAACACTTCAACAATAC
5	Left primer Right primer		197 482	25 25	56.79 59.79	56.00 44.00	9 4	TTTTTGTAAATTTAGGTTTTGGTTT CTCTTCCAAAATCAACCAATAAAAA

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

Product size: 286, Tm: 69.8, CpGs in product: 5

12/8/13

::   ::   :     :: ::      :  :  :
<pre>61 CCGACCAGGGGGTTGCGGTGATGCAGGGGCTGGGTCTTTGCAGGATACACAGTCACCTTA</pre>
<pre>121 GGATGGACTAGGAGAAAAAAAGGTAGAGAAGAATGAATCAGGAAGTTAGAGTCTCGTTGTT</pre>
<pre>181 CAGCTGTTTGTATGCTTCTCTGTAAACCCAGGCTCTGGCCTCGACCAGGCCTCCAGCACA :  :                        :::      ::  :    </pre>
<pre>241 GCTGGCCATACGCCCTCACAGTGTCATCGGCCTGGAATTTAATCGTGATAGTGTGGACCT</pre>
301       ATCAGATTTGAGAGATGTTATAAAAAATTTTATTTGTTTCTTCATAGCTTGAAATTGTCA           :
<pre>361 CGCATTGTTGAAGTGTTTACAAATCTCTGAAAGTACAGTGTGTATTAATTA</pre>
<ul> <li>421 CCTGAGCCAGGTTGCCTGGTTCAAATCCAAGGTCTGCCTTTTACTGGTTGATCCTGGAAG</li> <li>::::::::::::::::::::::::::::::::::::</li></ul>
481 AGTTTTTGATTCTTTTGTGT                     481 AGTTTTTGATTTTTTGTGT
**************************************
<pre>** * Upper row: Original sequence * * Lower row: Bisulfite modified sequence * * (For display, assume all CpG sites are methylated) * * ++ CpG sites * * :::: Non-CpG 'C' converted to 'T' * * &gt;&gt;&gt;&gt;&gt; Left primer * www.urogene.org/cgl-bin/methprimer_results.cgi</pre>

 12/8/13
 MethPrimer Results - MethPrimer - Li Lab, UCSF

 \*
 <<<<<<>Right primer

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 \*

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MethPrimer v1.1 beta

Li Lab, Department of Urology, UCSF

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

## MethPrimer result

Please cite MethPrimer: Li LC and Dahiya R. MethPrimer: designing primers for methylation PCRs. Bioinformatics. 2002 Nov;18(11):1427-31. PMID: 12424112 Percentage 40 60 80 88 0 400 bp 200 bp 600 bp 800 bp 1000 CpGII III R1 F1 R2 F2 💹 R3 📖 R4 F4 🖉 R.5 F5 Bisulfite PCR primer CpG Island Input Sequence MSP Primer Set Methylated-Specific 888 1333 Unmethylated-Specific 📼--Sequence Name: Sequence Length: 1001 CpG island prediction results (Criteria used: Island size > 100, GC Percent > 50.0, Obs/Exp > 0.6) 1 CpG island(s) were found in your sequence Size (Start - End) (393 - 791) Island 1 399 bp Primer picking results for bisulfite sequencing (or restriction) PCR GC% 'C's Sequence Start Size Tm Primer 110 25 54.83 52.00 9 AATTTTTGATTTTAAGTGATTTGTT 1 Left primer 7 239 27 54.57 51.85 AAACTATTTCTAAACCATTTTCTACTC Right primer Product size: 130, Tm: 67.3, CpGs in product: 5 110 25 54.83 52.00 9 AATTTTTGATTTTAAGTGATTTGTT 2 Left primer 238 26 53.27 53.85 7 AACTATTTCTAAACCATTTTCTACTC Right primer Product size: 129, Tm: 67.3, CpGs in product: 5 110 25 54.83 52.00 9 AATTTTTGATTTTAAGTGATTTGTT 3 Left primer 237 25 51.86 52.00 6 ACTATTTCTAAACCATTTTCTACTC Right primer Product size: 128, Tm: 67.4, CpGs in product: 5 
 57.24
 56.00
 6
 GATTGGGTTTTATTATGTTGGTTAG

 54.57
 51.85
 7
 AAACTATTTCTAAACCATTTTCTAAC
 75 25 4 Left primer Right primer 239 27 AAACTATTTCTAAACCATTTTCTACTC Product size: 165, Tm: 68.2, CpGs in product: 6 www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

ieun finier result

### MethPrimer Results - MethPrimer - Li Lab, UCSF

5 Left primer 110 25 54.83 52.00 9 AATTTTTGATTTTAAGTGATTTGTT Right primer 240 28 55.76 50.00 7 AAAACTATTTCTAAACCATTTTCTACTC Product size: 131, Tm: 67.2, CpGs in product: 5	
1 CCTCAGCCTCCCGAGCAGCTGGGACTACGGGTGTGCGCCACCACGCCCAGCTAATTTTTT :: : ::::++  :  :  :  ++    ++:: :: ++::  :	
<pre>61 GTATTTTTAGTAGAGACTGGGTTTCACCATGTTGGCCAGGCTGGTCTCGAACTCCTGACC                                    </pre>	
<pre>121 TCAAGTGATCTGCCCGACTCGGCCTCCCCAAAATGCTGGGATTCCAGGCGTGAGCCACAGC  :       : ::++ : ++ :::::      :     ::   ++   ::::! + 121 TTAAGTGATTTGTTCGATTCGGTTTTTTAAAATGTTGGGATTTTAGGCGTGAGTTATAGC &gt;</pre>	
<pre>181 GCCTGTCCTGCATGTTACTTTTGAATGAAACCGAGCAGAAATGGCCCAGAAACAGCCTT     +::   ::  :  :  :++  :      ::::    :  :</pre>	
<pre>241 GCATCCATCAAGGGGCACACGACCCCCCCCACTACCTCCCCCCTCAACCTTGGAAGATCATTT</pre>	
301 AACAAATTCTTTGGTTTGAACACTTGATGTTACCTTGCCACTGGGGATACATCCCTAACT   :     :            : :       :::   :	
<pre>361 CTAGACAGCAGGTTGTTAAACACGGGGCCTGGTATCCACTAGGCGTCCCATAAATGCTGC :    :  :           : ++   ::      : ::    ++ :::      </pre>	
<pre>421 CACTTTTGTGGTTCCGAGGAGGCGGCTCACTCCGTTCGGGGCTTGGCAGGAGTCGTGGAGT : :         :++     ++ :J: : :++  ++  :    :++      ++       421 TATTTTGTGGTTTCGAGGAGGCGGTTTATTTCGTTCGGGTTTGGTAGGAGTCGTGGAGT</pre>	
<pre>481 GGGTTCGGCCACGTGGAATCCGCGTCCTGGGAACCCGTGGAATCCGCGTCCTGGGAACCC      ++ :: ++     :++++ ::      ::+++     :++++!::      ::++ 481 GGGTTCGGTTACGTGGAATTCGCGTTTTGGGAATTCGTGGAATTCGCGTTTTGGGAATTC</pre>	
541 GTGGAATCCGCCTCCTGGGAACCCGTGGAATCCGCCTCCTGGGAACCCGTGGAATCCGCC +     :++:::::!!    ::++:::::!     :++:: 541 GTGGAATTCGTTTTTTGGGAATTCGTGGAATTCGTTTTTTGGGAATTCGTGGAATTCGTT	
601 TCCTGGGAACCCGTGGAATCCGCCTCCTGGGAACCCGTGGAATCCGCCTCCTGGGAACCC www.urogene.org/cgi-bin/methprimer/methprimer_results.cgi	

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/13		MethPrimer Results - MethPrimer - Li Lab, UCSF
601		::++      :++:: ::      ::++      :++:: ::
	+         [ +	CGGGTTGGATGCGCATGTGCGTGTCTCTTTTTCCGGGGGAGGCTCCGCCCAC +        ++:     ++   : :     :+++
	GCCCCGCC	CCCGCTCCCAAGTGCGCGCGGACCCCTAGCTCCCTCTGAGTTGCGCTGGGCT :++:!:::     ++++++  ::::!!!!::::!!      ++:  ]!!
		TCGTTTTTAAGTGCGCGCGCGGATTTTTAGTTTTTTTGAGTTGCGTTGGGTT GCACCATGACCCTGGAGGCGATCCGCTACTCGCGGGGCTCCCTGCAGATCCT/
	:  :	: ::    :::       ++  :++:  : ++++   : :::  :
	::  :	rGCTGCTGCCCAAGCAGAGCCGCTACGAGGCGGTGGGCTCGGTGCACCAGGC(   :  :  :::   :    :++:  ++   ++    : :++   : ::   : rGTTGTTGTTTAAGTAGAGTCGTTACGAGGCGGTGGGTTCGGTGTATTAGGT
		CCATCCGCGCCATGAAGGTGCAGCGGGGGGGGGGGGGGG
961	+   ++       - GGCGGGGG	CGGCGGGGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGG
· Ex	planation	ns
		Original sequence Bisulfite modified sequence (For display, assume all CpG sites are methylated)
k .		CpG sites
k K ++		~
€ * ++ * ::	::	Non-CpG 'C' converted to 'T'
++ + :: + >>		Non-CpG 'C' converted to 'T' Left primer Right primer

MethPrimer v1.1 beta

Li Lab, Department of Urology, UCSF

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

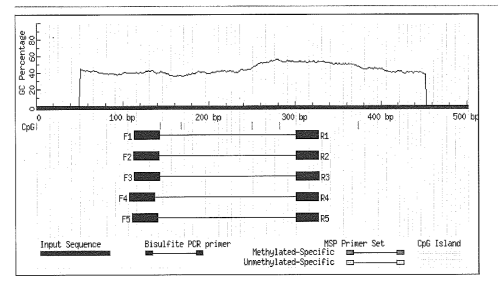
### MethPrimer Results - MethPrimer - Li Lab, UCSF

# **MethPrimer result**

Please cite MethPrimer: Li LC and Dahiya R. <u>MethPrimer: designing primers for methylation PCRs</u>. Bioinformatics. 2002 Nov;18(11):1427-31.

PMID: 12424112

12/8/13



Sequence Name: Sequence Length: 501

CpG island prediction results (Criteria used: Island size > 100, GC Percent > 50.0, Obs/Exp > 0.6) No CpG islands were found in your sequence Primer picking results for bisulfite sequencing (or restriction) PCR

1	Right primer	113 30 327 26	Tm GC% 'C' 56.56 33.33 4 59.23 65.38 10 CpGs in product: 4	TTGAGGAAAAATAATAAAGAGTTATTAGAT
2	Left primer Right primer Product size: 216,	327 26	56.56 33.33 4 59.23 65.38 10 CpGs in product: 4	АТТСАССАААААТААТАААСАСТТАТТАСА ААССТАААСААТАССАТТСААААСАА
3	Right primer	113 30 328 27 Tm: 64.0,		TTGAGGAAAAATAATAAAGAGTTATTAGAT TAACCTAAACAATACCATTCAAAACAA
4	Left primer Right primer Product size: 221,	327 26	55.61 30.00 4 59.23 65.38 10 CpGs in product: 4	ТАААТАТТGАGGAAAAATAATAAAGAGTTA ААССТАААСААТАССАТТСААААСАА
5	Left primer Right primer	111 30 327 26	55.34 33.33 4 59.23 65.38 10	TATTGAGGAAAAATAATAAAGAGTTATTAG AACCTAAACAATACCATTCAAAACAA

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

### MethPrimer Results - MethPrimer - Li Lab, UCSF

12/8/13

Product size: 217, Tm: 63.9, CpGs in product: 4

	:: :	TCACTTACCCAGGTTCAGGAAAGCATTTTCCAGGTCTCCTTTAACCTCTT  : :   :::    ::!!!! TTATTTATTTAGGTTTAGGAAAGTATTTTTTAGGTTTTTT	:
	:     :	TTCCAACATGTCATAAGGGCTGTAACTCTTGTACCTATCAAATACTGAGG   ::  :   :   :   :  :  :  :   ::   ::  :  :  : TTTTAATATGTTATAAGGGTTGTAATTTTTGTATTTATT	 AA
	:  :  AAATAATA	AAGAGTTATCAGATCCGAGCCACTAGTCAAAGCTGTCAACGATCACCCACG              ++  :: :    :    :  ++  : :::: : AAGAGTTATTAGATTCGAGTTATTAGTTAAAGTTGTTAACGATTATTAT >>>>>>>>>>>>>>>>>>>>>>>>>>>>>>	:
		"GCACCATAATTTTTTTAAAAATTGAGGATGATCACAGCATCCTAGGAGCT"   : ::	11
	:	CCACGGTGACCAGAGCCAACATTGGCCAAGTTTGTCGTGGAACAGCCATAC :: ++    ::    ::     ::     ++     :  :	:
	:: [ ::  TTTGTTTI	rGAATGGCACTGCCCAGGCCACATATTTGGACCATCTCTATCTCCCCTGAG        : : : :::   :::       ::  :  : :	11
	:::	TTCCATCCGAAAACCATAGGAAACAGTACAGAGCATGCACCAAAGTCCACT.   ::  :++    ::       :  !!!!!!!!!!!!!!	:
	[: ]: ]	AATAATGGCAAGACCAAATGATCATCAAACAAGAAGGAGCTGCAGAATAAA 	:
	::	GCAGAAACTATTTG  :    :  !   GTAGAAATTATTTG	
****	*******	*****	**
* Ex	planation	ns	*
* Up * Lc * * ++	per row: wer row:	Original sequence Bisulfite modified sequence (For display, assume all CpG sites are methylated) CpG sites Non-CpG 'C' converted to 'T' Left primer	_* * * * * *

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

 12/2/13
 MethPrimer Results - MethPrimer - Li Lab, UCSF

 \*
 <<<<<>Right primer

 \*
 \*

 \*
 \*

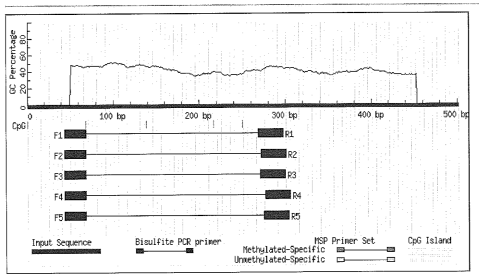
MethPrimer v1.1 beta

Li Lab, Department of Urology, UCSF

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

#### MethPrimer Results - MethPrimer - Li Lab, UCSF

#### MethPrimer result



Please cite MethPrimer: Li LC and Dahiya R. <u>MethPrimer: designing primers for methylation PCRs</u>. Bioinformatics. 2002 Nov;18(11):1427-31.

PMID: <u>12424112</u>

12/8/13

Sequence Name: Sequence Length: 501

CpG island prediction results (Criteria used: Island size > 100, GC Percent > 50.0, Obs/Exp > 0.6) No CpG islands were found in your sequence Primer picking results for bisulfite sequencing (or restriction) PCR

1	Right primer	44 2·5 298 30	Tm GC% 56.36 56.00 55.74 36.67 CpGs in product	7 5	-
2	Left primer Right primer Product size:		56.36 56.00 55.41 36.67 CpGs in product	5	TTTAGGTGTGTGTGTTTATTTTTAGGA CAATAACCTTATCTAATTTATCATTTTAAC
3	Left primer Right primer Product size:	300 30	56.36 56.00 55.41 33.33 CpGs in product	5	TTTAGGTGTGTGTGTTTATTTTTAGGA AATAACCTTATCTAATTTATCATTTTAACA
4	5 1	306 30	56.36 56.00 56.35 43.33 CpGs in product	8	TTTAGGTGTGTGTGTTTATTTTTAGGA AAAAACAATAACCTTATCTAATTTATCATT
5	Left primer Right primer	44 25 304 30	56.36 56.00 56.35 43.33	7 8	TTTAGGTGTGTGTGTTTATTTTTAGGA AAACAATAACCTTATCTAATTTATCATTTT

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

12/8/13

Product size: 261, Tm: 68.8, CpGs in product: 4

	AGTGTAGGAGAGGTGAGTTGCTTAGGTCTAAGGAGAAAGACTGCTTAGGTGTGTGT
	CCCCAGGACGAAGGAAGGAACACTGGGTGAGATTTTGTTCAACTACCCATAGTTACCACC ::::    ++
	AGATGGTGAAACTGATCCCGGGCCTCTTGGGTATTGATCAGTTTATGGGGAGATGGGGAG 
	AAGACTATCTTTCACTTGTTAATTCATTAATTTCTTTCGCAAATATTTTTTCAGTACCTG     :  :  :  :  :  :     :  :   :++:!      :   :
	CTAAGTCCCACGGACTATGCTAGGAGCTGCTGTTAAAATGACAAACCAGATAAGGTCACT :     ::: ++  :    :  :  :  :       : ::   :    : :: TTAAGTTTTACGGATTATGTTAGGAGTTGTTGTTGTTAAAATGATAAATTAGATAAGGTTATT <<<<<<<<<<
	GCCCTTAATCAACTTACAGTTGGGTGAGAAGCTATCAGGTACAAGTATGGCCCTAGAACA  ::::     :  :  :  :   :   :  :  :   :  :
	AATTAGTCTTTTCTAGTTAATAATCTTATGTGATGAGATTTGGCCTTGCTCCTTTGGTGA
	CTTGCCTCAAGGAGCCCCAGGCAAAACCAATGTAACATATATTAATAATATATGAAATAA :\\\::\:\:\\\\\\\:::\\\\\\\\::\\\\\\\\
	TATATTTGTAGACACAATTG              :!:      TATATTTTGTAGATATAATTG
	**************************************
* Lc * * ++ * ::	per row: Original sequence * wer row: Bisulfite modified sequence * (For display, assume all CpG sites are methylated) *

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

12/8/13 MethPrimer Results - MethPrimer - Li Lab, UCSF \* <<<<< Right primer \* 

\*

\*

MethPrimer v1.1 beta

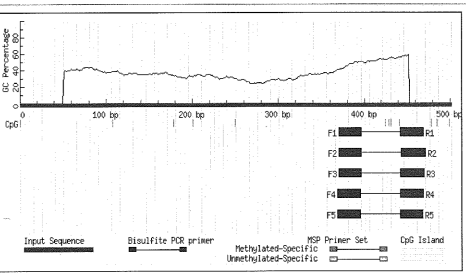
Li Lab, Department of Urology, UCSF

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www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

#### MethPrimer Results - MethPrimer - Li Lab, UCSF

#### **MethPrimer result**



Please cite MethPrimer: Li LC and Dahiya R. <u>MethPrimer: designing primers for methylation PCRs</u>. Bioinformatics. 2002 Nov;18(11):1427-31.

PMID: <u>12424112</u>

12/8/13

Sequence Name: Sequence Length: 501

CpG island prediction results (Criteria used: Island size > 100, GC Percent > 50.0, Obs/Exp > 0.6) No CpG islands were found in your sequence Primer picking results for bisulfite sequencing (or restriction) PCR

	Primer	S	tart	Size	Tm	GC%	'C's	Sequence
1	Left primer							1
_								TCCTCCTACAATACTAAAATTACAAAC
	Product size: 1							
2	Left primer		370	26	59.41	50.00	4	TAGGGGAAAATTGAGGTTATAAGAAG
	Right primer .		471	29	58.29	55.17	7	CATCCTCCTACAATACTAAAATTACAAAC
	Product size: 3	102,	Tm:	60.0,	CpGs in	product	: 4	
3	Left primer		370	26	59.41	50.00	4	TAGGGGAAAATTGAGGTTATAAGAAG
	Right primer		470	28	56.18	53.57	7	ATCCTCCTACAATACTAAAATTACAAAC
	Product size: 3	101,	Tm:	60.1,	CpGs in	product	: 4	
4	L.				59.64			ATAGGGGAAAATTGAGGTTATAAGAAG
	Right primer		469	27	55.81	55.56	7	TCCTCCTACAATACTAAAATTACAAAC
	Product size: 3	101,	Tm:	60.1,	CpGs in	product	: 4	
5	Loft primor		369	27	59.64	48.15	4	ATAGGGGAAAATTGAGGTTATAAGAAG
5	Left primer							
	Right primer		468	26	54.24	57.69	7	CCTCCTACAATACTAAAATTACAAAC

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

Product size: 100, Tm: 60.1, CpGs in product: 4

1 ATCATATATGAACTTCTGTTTTTGAAGCCACCCCTCAAGAGCCAACAGGATTCTTTAAGT           :                    ::::::::::::::
61 ATCCCAGTGGTACTAAACCCATATCCTTTGAGAATGCTTTCTTAGAACGATTCACAAACT   :::       :   ::    ::    :
121 GGCTCTGCAGGCTTTTCAAAACTTAAGTTCTAGAAGTTGTACAATAAAATGACAGGATCG   : : :  :  :  :   :   :   :    :
181 CTAAAATAAGTGTATGGCATTCGATGTAACTGCTTGCAAAAACAACATCCAATTTTAATA :
241 TTGGCCTAATCGTGGCTAAATATTGGTATAATAATAGTTAACTATCTGCTAAGTTCTATT     :::   ++   :
301 TTAAAGCTTTATTTATTTATTATCACACAACAAACCTGTAAGGTGGGTCTCATTAGCAGTCT       :               : :  :   :
<pre>361 CATTTTACACAGGGGAAAACTGAGGCTACAAGAAGTAACTTGTTAAAGGTTATGCAGCTA :      : :       :   :  :  :  :  :  :  </pre>
421 GAGGCCGGGCGCGGTGGCTCACGCCTGTAATCCCAGCACTGTAGGAGGATGAGGCAGACG     :+++  ++++   :::++::     :::  :  :
<pre>481 GATCACGAGGTCAGGAGATCG</pre>
<pre>************************************</pre>

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

 12/8/13
 MethPrimer Results - MethPrimer - Li Lab, UCSF

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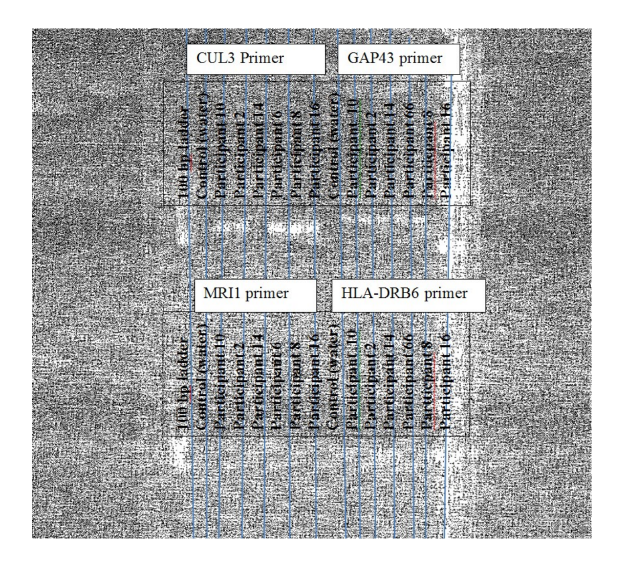
 \*
 \*

MethPrimer v1.1 beta

Li Lab, Department of Urology, UCSF

www.urogene.org/cgi-bin/methprimer/methprimer\_results.cgi

Appendix F PCR product Images



100bpLadder Control(water) Participant10 Participant2 Participant8 Participant8 Participant16 Control(water) Participant10 Participant14 Participant14 Participant16 Participant16 Participant16 Participant16 Participant16	100bpLadder Control(water) Participant10 Participant2 Participant6 Participant6 Participant16 Control(water) Participant16 Participant14 Participant14 Participant6 Participant6 Participant6 Participant6 Participant6 Participant16 Participant16	ANXA	A2 Prim er	RHPN1 P	Prim er	1
		100bpLadder Control(water) Participant10 Participant2	Participant14 Participant6 Participant8 Participant16	Control (water) Participant 10 Participant 2 Participant 14	Participant6 Participant8 Participant16	
						Colle Black

Appendix G

Samples of Inadequate Quality for Sequencing

RHPN1

```
CGAGGCGCAGCCTGGTGCGGGCGCCACGGGGTCGGGCTGTGATCGCCTGTGG
CCTCCCTGCAGGGCTGTGACTCCCTGACGCAGATCCAGTGCGGCCAGCTGCA
GAGCCGCAGGGCCCAGATTCACCAGCAGATTGACAAGGAGCTGCAGATGCG
GACGGGCGCTGAGAACCTCTACAGGTCAGTGCTTGAGACTGCCCGGCCCCGG
```

excluded: RHPN16216FWD-RHPN1FWD Sequence identity: 89.2% excluded: RHPN16216REV-RHPN1REV Sequence identity: 89.2% excluded: RHPN16217FWD-RHPN1REV Sequence identity: 87.2% excluded: RHPN16228FWD-RHPN1FWD Sequence identity: 87.7% excluded: RHPN16228REV-RHPN1REV Sequence identity: 89.5% excluded: RHPN16235FWD-RHPN1FWD Sequence identity: 88.6% excluded: RHPN16235REV-RHPN1REV Sequence identity: 15.4% excluded: RHPN16252FWD-RHPN1FWD Sequence identity: 88.2% excluded: RHPN16259FWD-RHPN1FWD Sequence identity: 76.6% excluded: RHPN16259FWD-RHPN1FWD Sequence identity: 86.3% excluded: RHPN16217FWD-RHPN1FWD Sequence identity: 90.2%; Conversion rate: 100.0%; N-sites at non CpG cytosines positions: 5.9%; N-sites at CpG positions: 28.6%; Gaps: 1.4%

1 out of <u>12 uploaded</u> sequencing results pass the quality criteria when compared to the reference sequence. At the next step all sequences are compared against all others to detect clonal amplifications as <u>described in the manual</u>.

# Sequence Alignment of the sequences included:

# RHPN16252REV-RHPN1RE

# ANXA

# CCGAGCCACTAGTCAAAGCTGTCAACGATCACCCACCTAGTTTTATGCACCAT AATTTTTTAAAAATTGAGGATGATCACAGCATCCTAGGAGCTTAGAGGTTAC CACGGTGACCAGAGCCAACATTGGCCAAGTTTGTCGTGGAACAGCCATACCA C

```
excluded: ANXA26216FWD-ANXA2FWD Sequence identity: 65.0%
excluded: ANXA26216FWD-ANXA2FWD_R Sequence identity: 71.5%
excluded: ANXA26216REV-ANXA2REV Sequence identity: 80.9%
excluded: ANXA26216REV-ANXA2REV_R Sequence identity: 64.6%
excluded: ANXA26217FWD-ANXA2FWD Sequence identity: 68.8%
excluded: ANXA26217FWD-ANXA2REV Sequence identity: 79.7%
excluded: ANXA26228FWD-ANXA2FWD Sequence identity: 67.1%
excluded: ANXA26228REV-ANXA2REV Sequence identity: 69.7%
excluded: ANXA26235FWD-ANXA2FWD Sequence identity: 78.5%
excluded: ANXA26235REV-ANXA2REV Sequence identity: 67.7%
```

excluded: ANXA26252FWD-ANXA2FWD Sequence identity: 65.6% excluded: ANXA26252REV-ANXA2REV Sequence identity: 79.6% excluded: ANXA26259FWD-ANXA2FWD Sequence identity: 57.0% excluded: ANXA26259REV-ANXA2REV Sequence identity: 51.3%

No sequence passed the chosen sequence identity threshold of 90%

# HLA-DRB6

AAGGTAGAGAGAATGAATCAGGAAGTTAGAGTCTCGTTGTCAGCTGTTTGTA TGCTTCTCTGTAAACCCAGGCTCTGGCCTCGACCAGGCCTCCAGCACAGCTGG CCATACGCCCTCACAGTGTCATCGGCCTGGAATTTAATCGTGATAGTGTGGAC CTATCAGATTTGAGAGATGTTATAAAAAATTTTATTTGTTTCTTCATAGCTTG AAATTGTCAC

excluded: HLADBR66216FWD-HLADBR6FWD Sequence identity: 36.5% excluded: HLADBR66216REV-HLADBR6REV Sequence identity: 23.4% excluded: HLADBR66217FWD-HLADBR6FWD Sequence identity: 18.3% excluded: HLADBR66217REV-HLADBR6REV Sequence identity: 18.0% excluded: HLADBR66228FWD-HLADBR6FWD Sequence identity: 13.6% excluded: HLADBR66228REV-HLADBR6REV Sequence identity: 26.3% excluded: HLADBR66235FWD-HLADBR6FWD Sequence identity: 17.2% excluded: HLADBR66235FWD-HLADBR6FWD Sequence identity: 13.2% excluded: HLADBR66252FWD-HLADBR6FWD Sequence identity: 44.2% excluded: HLADBR66252FWD-HLADBR6FWD Sequence identity: 33.3% excluded: HLADBR66259FWD-HLADBR6FWD Sequence identity: 33.3% excluded: HLADBR66259FWD-HLADBR6FWD Sequence identity: 27.2%

# No sequence passed the chosen sequence identity threshold of 90%

# MRI1

# CGACTCGGCCTCCCAAAATGCTGGGATTCCAGGCGTGAGCCACAGCGCCTGT CCTGCATGTTACTTTTGAATGAAACC

excluded: MRI16216FWD-MRI1FWD Sequence identity: 37.8% excluded: MRI16216FWD-MRI1FWD\_R Sequence identity: 40.3% excluded: MRI16216REV-MRI1REV Sequence identity: 37.7% excluded: MRI16216REV-MRI1REV\_R Sequence identity: 55.8% excluded: MRI16217FWD-MRI1FWD Sequence identity: 33.8% excluded: MRI16217REV-MRI1REV Sequence identity: 44.2% excluded: MRI16228FWD-MRI1FWD Sequence identity: 51.9% excluded: MRI16228FWD-MRI1FWD Sequence identity: 64.4% excluded: MRI16235FWD-MRI1FWD Sequence identity: 50.6% excluded: MRI16235FWD-MRI1REV Sequence identity: 35.1% excluded: MRI16252FWD-MRI1FWD Sequence identity: 30.6% excluded: MRI16252REV-MRI1REV Sequence identity: 36.0% excluded: MRI16259FWD-MRI1FWD Sequence identity: 44.6% excluded: MRI16259REV-MRI1REV Sequence identity: 68.4%

No sequence passed the chosen sequence identity threshold of 90%

GAP43

excluded: GAP436216FWD-GAP43FWD Sequence identity: 54.8% excluded: GAP436216REV-GAP43REV Sequence identity: 44.3% excluded: GAP436217FWD-GAP43FWD Sequence identity: 70.1% excluded: GAP436217REV-GAP43REV Sequence identity: 31.2% excluded: GAP436228FWD-GAP43FWD Sequence identity: 74.6% excluded: GAP436228REV-GAP43REV Sequence identity: 74.6% excluded: GAP436235FWD-GAP43FWD Sequence identity: 76.6% excluded: GAP436235FWD-GAP43FWD Sequence identity: 36.0% excluded: GAP436252FWD-GAP43FWD Sequence identity: 80.3% excluded: GAP436252FWD-GAP43FWD Sequence identity: 48.9% excluded: GAP436259FWD-GAP43FWD Sequence identity: 60.7% excluded: GAP436259FWD-GAP43REV Sequence identity: 60.7%

#### No sequence passed the chosen sequence identity threshold of 90%

CUL3

#### TAACTTGTTAAAGGTTATGCAGCTAGAGGCCGGGGCGCGGGGGGCGCGCGGCGCCAC

excluded: CUL36216FWD-CUL3FWD Sequence identity: 45.7% excluded: CUL36216REV-CUL3REV Sequence identity: 60.9% excluded: CUL36217FWD-CUL3FWD Sequence identity: 56.5% excluded: CUL36217REV-CUL3REV Sequence identity: 54.3% excluded: CUL36228FWD-CUL3FWD Sequence identity: 50.0% excluded: CUL36228REV-CUL3REV Sequence identity: 60.9% excluded: CUL36235FWD-CUL3FWD Sequence identity: 45.7% excluded: CUL36235REV-CUL3REV Sequence identity: 52.2% excluded: CUL36252FWD-CUL3FWD Sequence identity: 54.3% excluded: CUL36252FWD-CUL3FWD Sequence identity: 47.8% excluded: CUL36259FWD-CUL3FWD Sequence identity: 54.3% excluded: CUL36259FWD-CUL3FWD Sequence identity: 54.3% excluded: CUL36259FWD-CUL3FWD Sequence identity: 54.3%

#### No sequence passed the chosen sequence identity threshold of 90%

Appendix I

R Code and Normalization Figures and Output Table

```
library(BiocInstaller)
biocValid()
source("http://bioconductor.org/biocLite.R")
biocLite("limma")
biocLite("illuminaio")
biocLite("minfi")
biocLite("minfiData")
biocLite() ##installs every package on bioconductor
biocLite("IlluminaHumanMethylation450kmanifest")
require("minfi")
require("minfiData")
##generate a Red Green data set
baseDir<-("C:\\Users\\Michelle\\Dropbox\\dissertation\\iDat all")
targets <- read.450k.sheet(baseDir)
RGset<-read.450k.exp(base=baseDir, targets=targets)
pd<-pData(RGset)
gcReport(RGset, sampNames = pd$Sample ID,sampGroups=pd$GBS,pdf =
"qcReport.pdf")
densityPlot(RGset, sampGroups = pd$GBS,main = "Beta", xlab = "Beta")##shows
density plot in R workspace
par(oma=c(2,10,1,1))
densityBeanPlot(RGset, sampGroups = pd$GBS, sampNames = pd$Sample ID)
##Methylation sets
MSet.raw <- preprocessRaw(RGset)
MSet.norm <- preprocessIllumina(RGset, bg.correct = TRUE, normalize = "controls",
reference = 2) ##normalizes methylation data
controlStripPlot(RGset, controls="BISULFITE CONVERSION II", sampNames =
pd$Sample ID)##control bisulfite conversion II
annot<-read.csv("http://supportres.illumina.com/documents/myillumina/b78d361a-def5-
4adb-ab38-e8990625f053/humanmethylation450 15017482 v1-2.csv",
skip=7,head=TRUE)
getMeth(MSet.raw)[1:4,1:3]##returns raw methylation values use
getUnmeth(MSet.raw)[1:4,1:3]##returns raw unmethylated values
getBeta(MSet.raw, type = "Illumina")[1:4,1:3] ##returns raw beta values
MSet.raw<-preprocessRaw(RGset)
Beta.raw<-getBeta(MSet.raw)
rownames(Beta.raw)[1:5]
annot$IlmnID[1:5]
annot<-annot[match(rownames(Beta.raw),annot$IlmnID),,drop=FALSE] ##annotates
betafile with illumina IDs
class(rownames(Beta.raw))
class(annot$IlmnID)
all.equal(as.character(annot$IlmnID),rownames(Beta.raw))
names(annot)
head(annot)
```

```
table(annot$Probe SNPs)
##first removed CpG sites that were known to be associated with SNP
any.SNPs<-ifelse(annot$Probe SNPs!="",1,0)
raw.wo<-Beta.raw[any.SNPs!=1,]
dim(raw.wo)
annot.wo<-annot[any.SNPs!=1,]
dim(annot.wo)
all.equal(as.character(annot.wo$IlmnID),rownames(raw.wo))
sum(raw.wo==1,na.rm=TRUE)
sum(raw.wo==0,na.rm=TRUE)
## To avoid errors when applying the logit transformation that would occur if beta=0 or
beta=1.
## we imputed 0.999 when beta=1 and 0.001 when beta=0
raw.wo[raw.wo==1]<-0.999
raw.wo[raw.wo==0]<-0.001
sum(raw.wo==1,na.rm=TRUE)
sum(raw.wo==0,na.rm=TRUE)
#After imputation, the logit transformation was applied to the beta values
logit<-log(raw.wo/(1-raw.wo)) ## Logit transformed beta values
hist(logit[1,])
hist(raw.wo[1,])
Peak.correction<-
function(exprs,annot) {
     for (i in 1:dim(exprs)[2]) {
    dens.I<-density(exprs[annot$Infinium Design Type=="I", i],na.rm=T)
     dens.II<-density(exprs[annot$Infinium Design Type=="II", i],na.rm=T)
     sigma.uII<- -dens.II$x[dens.II$x<0][which.max(dens.II$v[dens.II$x<0])]
     sigma.mII<-dens.II$x[dens.II$x>0][which.max(dens.II$y[dens.II$x>0])]
     sigma.uI<- -dens.I$x[dens.I$x<0][which.max(dens.I$v[dens.I$x<0])]
     sigma.mI<-dens.I$x[dens.I$x>0][which.max(dens.I$y[dens.I$x>0])]
     exprs[,i]<- ifelse(annot$Infinium Design Type=="II" & exprs[,i]<0,
       exprs[,i]/sigma.uII*sigma.uI,
      ifelse(annot$Infinium Design Type=="II" & exprs[,i]>0,
            exprs[,i]/sigma.mII*sigma.mI, exprs[,i]))
}
     exprs
```

all.equal(as.character(annot.wo\$IlmnID),rownames(logit) ##The logit transformed beta values were then peak corrected to ensure the peaks of the Infinium II

##design beadtype were comparable to the peak locations of the Infinum I beadtype correct.methyl<-Peak.correction(logit,annot.wo)

}

pData(MSet.raw)

plot(density(logit[annot.wo\$Infinium\_Design\_Type=="I",1]), col="black", xlab="M-values",main="Sample 1", ylim=c(0,0.3))

lines(density(logit[annot.wo\$Infinium\_Design\_Type=="II",1]), col="red")

lines(density(logit[annot.wo\$Infinium\_Design\_Type=="II",1],na.rm=TRUE), col="red") plot(density(correct.methyl[annot.wo\$Infinium\_Design\_Type=="I",1]), col="black",

xlab="M-values",main="Peak-corrected Sample 1", ylim=c(0,0.3))

lines(density(correct.methyl[annot.wo\$Infinium\_Design\_Type=="II",1],na.rm=TRUE), col="red")

plot(density(logit[annot.wo\$Infinium\_Design\_Type=="I",4]), col="black", xlab="M-values",main="Sample 4", ylim=c(0,0.3))

lines(density(logit[annot.wo\$Infinium\_Design\_Type=="II",4],na.rm=TRUE), col="red") plot(density(correct.methyl[annot.wo\$Infinium\_Design\_Type=="I",4]), col="black", xlab="M-values",main="Peak-corrected Sample 4", ylim=c(0,0.3))

lines(density(correct.methyl[annot.wo\$Infinium\_Design\_Type=="II",4],na.rm=TRUE), col="red")

plot(density(logit[annot.wo\$Infinium\_Design\_Type=="I",7]), col="black", xlab="M-values",main="Sample Unmethy", ylim=c(0,0.3))

lines(density(logit[annot.wo\$Infinium\_Design\_Type=="II",7],na.rm=TRUE), col="red") plot(density(correct.methyl[annot.wo\$Infinium\_Design\_Type=="I",7]), col="black", xlab="M-values",main="Peak-corrected Sample unmeth", ylim=c(0,0.3))

lines(density(correct.methyl[annot.wo\$Infinium\_Design\_Type=="II",7],na.rm=TRUE), col="red")

##this is unnecessary because GBS is either positive or negative so a subset does not need to be created;

##however other data sets may need subsets created to carry out analysis and here for your reference

```
Msubset<-correct.methyl[,pData(MSet.raw)$GBS=="Negative"|
```

pData(MSet.raw)\$GBS=="Positive"]

##Notice results for GBS will be the same as the above using

```
Msubset2<-correct.methyl
```

```
all.equal(Msubset,Msubset2)
```

```
dim(Msubset)
```

class(Msubset)

```
##if you need a group vector, this is how it should be formatted
group<-pData(MSet.raw)[pData(MSet.raw)$GBS=="Negative"]
```

```
pData(MSet.raw)$GBS=="Positive","GBS"]
```

##However, I don't need to use one because of how my data is structured; ##instead I could use group<-pData(MSet.raw)\$GBS as the vector in design pData(MSet.raw)

library(limma)

design<-model.matrix(~as.factor(group)-1)

design

colnames(design)<-c("Negative","Positive")

design

```
fit<-lmFit(Msubset, design) ##least squares fitting off linear model for each gene
fit
contr.matrix<-makeContrasts(GBS.status=Positive-Negative, levels=design)
contr.matrix
fit2<-contrasts.fit(fit, contr.matrix) #makes contras to compare by GBS status
fit3<-eBayes(fit2) ##use of array weights increases the significance of top genes
results<-topTable(fit3, coef="GBS.status", number=dim(Msubset)[1],
 sort.by="none", adjust="BH")
### Warning, the sort.by="none" and number=dim()[1] parameters are important to
ensure your results are aligned with the annotation data
head(results)
results<-data.frame(ID=rownames(results),results)
head(results)
head(Msubset)
dim(results)
dim(Msubset)
all.equal(results$ID, rownames(Msubset$ID))
class(results$ID)
class(Msubset)
class(Msubset[,1])
all.equal(comp[,1],comp[,2])
all.equal(results$ID,Msubset[,1])
### Note that without specifying sort.by="none" the annotation information is misaligned
##correct.beta is the back-transformed peak corrected logitvalues
correct.beta<-exp(Msubset)/(1+exp(Msubset))
dim(correct.beta)
head(correct.beta)
sum(results$adj.P.Val<0.05)
mu.Negative<-apply(correct.beta[,group=="Negative"], 1, mean)
mu.Positive<-apply(correct.beta[,group=="Positive"], 1, mean)
delta.beta<-mu.Positive-mu.Negative
mu.Negative[1:10]
mu.Positive[1:10]
delta.beta[1:10]
all.equal(results$ID.as.character(annot.wo$IlmnID))
class(results$ID)
class(annot.wo$IlmnID)
```

```
class(annot.wo$IImnID)
all.equal(comp[,1],comp[,2])
all.equal(results$ID,annot.wo$IImnID)
class(results$ID)==class(annot.wo$IImnID)
head(annot.wo)
head(results)
results<-data.frame(ID=rownames(results),results)
```

### Note that final.results and annot.wo are matched so we can append annot.wo to final results;

## using generally accepted FDR cut off standards final.FDR.001results<-data.frame(Probe=rownames(results), beta.Negative=mu.Negative, beta.Positive=mu.Positive, delta.beta=delta.beta, p.value=results\$P.Value, FDR=results\$adj.P.Val, annot.wo) sign.results<-final.results[final.results\$FDR<0.001 & abs(final.results\$delta.beta)>0.2,] dim(sign.FDR.001results) # yields 0 rows = zero significantly different CpG sites final.FDR.05results<-data.frame(Probe=rownames(results), beta.Negative=mu.Negative, beta.Positive=mu.Positive, delta.beta=delta.beta, p.value=results\$P.Value, FDR=results\$adj.P.Val, annot.wo) sign.results<-final.results[final.results\$FDR<0.05 & abs(final.results\$delta.beta)>0.2,] dim(sign.FDR.05results) # yields 0 rows = zerp significantly differnt CpG sites final.FDR.01results<-data.frame(Probe=rownames(results), beta.Negative=mu.Negative, beta.Positive=mu.Positive, delta.beta=delta.beta, p.value=results\$P.Value, FDR=results\$adj.P.Val, annot.wo) sign.results<-final.results[final.results\$FDR<0.01 & abs(final.results\$delta.beta)>0.2,] dim(sign.results) # yields 0 rows = zero significantly differnt CpG sites ##Volcano plot, showing no significant differences. Signifincat CpG sites would be blue install.packages("ggplot2") require(ggplot2) ##Highlight CpGs that have an absolute fold change > 2 and a FDR < 0.05FDRcut = as.factor(abs(results logFC) > 2 & results adj.P.Val < 0.05)sum(abs(results logFC) > 2 & results adj.P.Val < 0.05)##Construct the plot object g = ggplot(data=results, aes(x=logFC, y=-log10(P.Value), colour= FDRcut)) + geom point(alpha=0.4, size=1.75) +

theme(legend.position = "none") + xlim(c(-4, 4)) + ylim(c(0, 6)) + xlab("log2 fold change") + ylab("-log10 p-value") ##returns no blue (significant) dots, all pink CpGs

### using generic p-value 0.05 cutoff that Joyce uses final.results<-data.frame(Probe=results\$ID, beta.Negative=mu.Negative, beta.Positive=mu.Positive, delta.beta=delta.beta, p.value=results\$P.Value, FDR=results\$adj.P.Val, annot.wo) sign.results<-final.results[final.results\$p.value<.05& abs(final.results\$delta.beta)>0.2,] dim ### To output in order to email your findings to an investigator, use write.table write.table(sign.results,"significantRrun.csv",sep=",",row.names=FALSE) ### You could also remove some of the annotation fields using -c() in square bracket notation; investigators may not be interested in all fields. ### bisulfitesequencing validation code

getBeta(MSet.raw)["cg11164659",] ##RHPN1 raw uncorrected beta value getBeta(MSet.raw)["cg09785377",] ##ANXA2 raw uncorrected beta value

cpgsD<-c("cg09785377","cg11164659")

cpgsD

plotCpg(Msubset, cpg=cpgsD[1], pheno=pData(Msubset)\$GBS) #ANXA2 plot of raw beta values by GBS status for each participant plotCpg(Msubset, cpg=cpgsD[2], pheno=pData(Msubset)\$GBS) ##RHPN1 plot of raw

beta values by GBS status for each participant

(correct.beta)["cg11164659",] ##RHPN1 normalized beta value (correct.beta)["cg09785377",] ##ANXA2 normalized beta value

##raw and corrected beats for CpG site only significant using R platform compared to genome studio significant values getBeta(MSet.raw)["cg23947138",] ##RASA3 raw beta value

(correct.beta)["cg23947138",] ##RASA3 normalized beta value

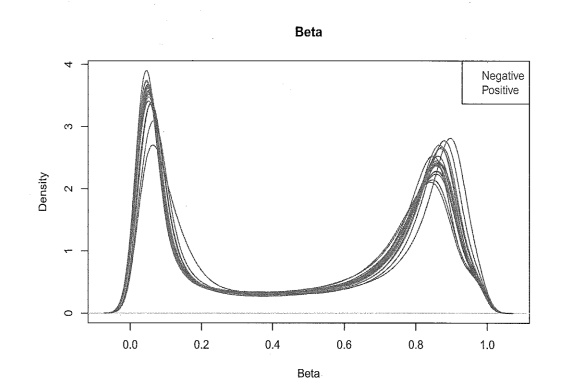
getBeta(MSet.raw)["cg00540295",] ##FAM69B raw beta value

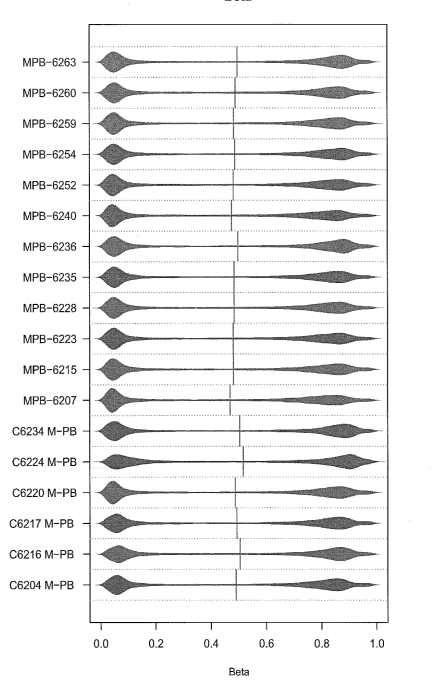
(correct.beta)["cg00540295",] ##FAM69B - snp

getBeta(MSet.raw)["cg01270299",] ##ZNF137 - snp (correct.beta)["cg01270299",]

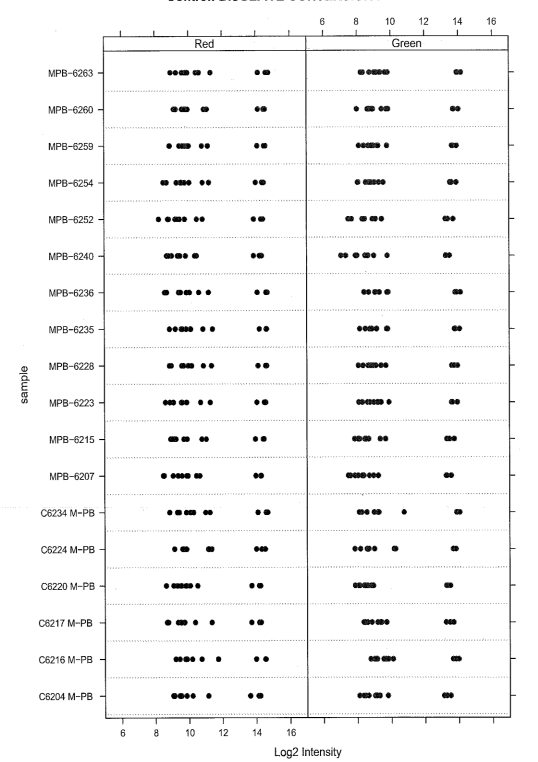
getBeta(MSet.raw)["cg25909532",] ##VIPR2 - snp (correct.beta)["cg25909532",]

getBeta(MSet.raw)["cg06688803",] ##CLPTM1 - snp (correct.beta)["cg06688803",] getBeta(MSet.raw)["cg07304760",] ##SND1 - reads (correct.beta)["cg07304760",] getBeta(MSet.raw)["cg15290312",] ##TIMP2 - snp (correct.beta)["cg15290312",] getBeta(MSet.raw)["cg10058204",] ##FLJ37201 - reads (correct.beta)["cg10058204",] getBeta(MSet.raw)["cg05331763",] ##FOxK2 - reads (correct.beta)["cg05331763",] getBeta(MSet.raw)["cg12434901",] ##KCNH6 - reads (correct.beta)["cg12434901",] getBeta(MSet.raw)["cg24634471",] ##JRK - reads (correct.beta)["cg0421902",] ##ZNF665 - reads (correct.beta)["cg01421902",] getBeta(MSet.raw)["cg04388792",] ##ZNF490 - reads (correct.beta)["cg04388792",] getBeta(MSet.raw)["cg13506281",] ##MTUS2 - snp (correct.beta)["cg04388792",] getBeta(MSet.raw)["cg07703391",] ##BMP8B/PPIE -reads (correct.beta)["cg07703391",] getBeta(MSet.raw)["cg10632770",] ##KIAA1199 -snp (correct.beta)["cg10632770",] getBeta(MSet.raw)["cg20479209",] ##FLJ43860 -reads (correct.beta)["cg20479209",] getBeta(MSet.raw)["cg13066461",] ##MRGPRX2-reads (correct.beta)["cg13066461",] getBeta(MSet.raw)["cg21130926",] ##SULF2 -reads (correct.beta)["cg21130926",] getBeta(MSet.raw)["cg04922606",] ##FAM120B -SNP (correct.beta)["cg04922606",] getBeta(MSet.raw)["cg03292225",] ##TNNT3 -reads (correct.beta)["cg03292225",] getBeta(MSet.raw)["cg10890644",] ##TUBAL3- snp (correct.beta)["cg10890644",] getBeta(MSet.raw)["cg17671604",] ##SPTBN4- snp (correct.beta)["cg17671604",] getBeta(MSet.raw)["cg09307883",] ##ANAPC2- reads (correct.beta)["cg09307883",] getBeta(MSet.raw)["cg10528424",] ##SYT8 - reads (correct.beta)["cg10528424",] getBeta(MSet.raw)["cg07480176",] ##CASD1 - snp (correct.beta)["cg07480176",] getBeta(MSet.raw)["cg14252149",] ##LGALS8 -snp (correct.beta)["cg14252149",] getBeta(MSet.raw)["cg12155450",] ##NAT14/ZNF628 -reads (correct.beta)["cg12155450",] getBeta(MSet.raw)["cg20976286",] ##OCA2 - reads (correct.beta)["cg20976286",] getBeta(MSet.raw)["cg10995422",] ##HLA-DRB6 - snp (correct.beta)["cg10995422",]

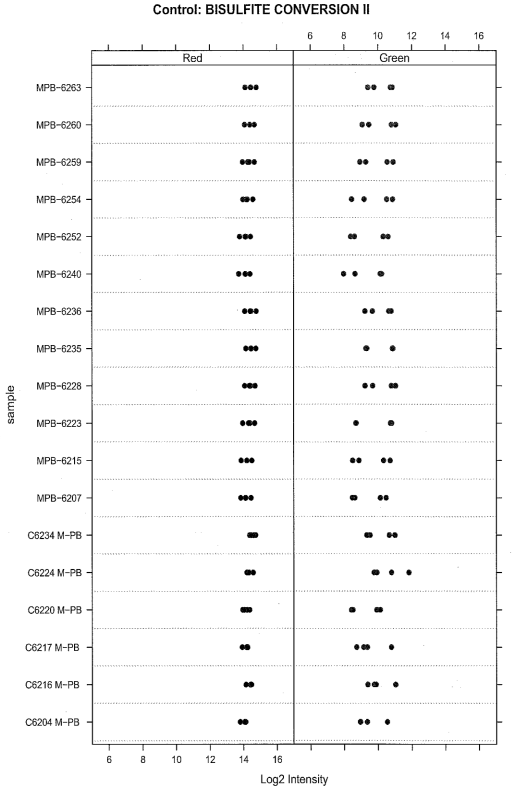




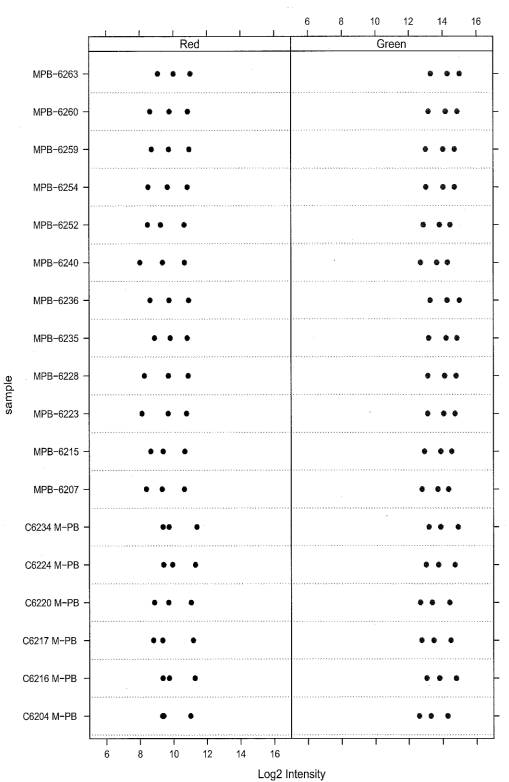
Beta



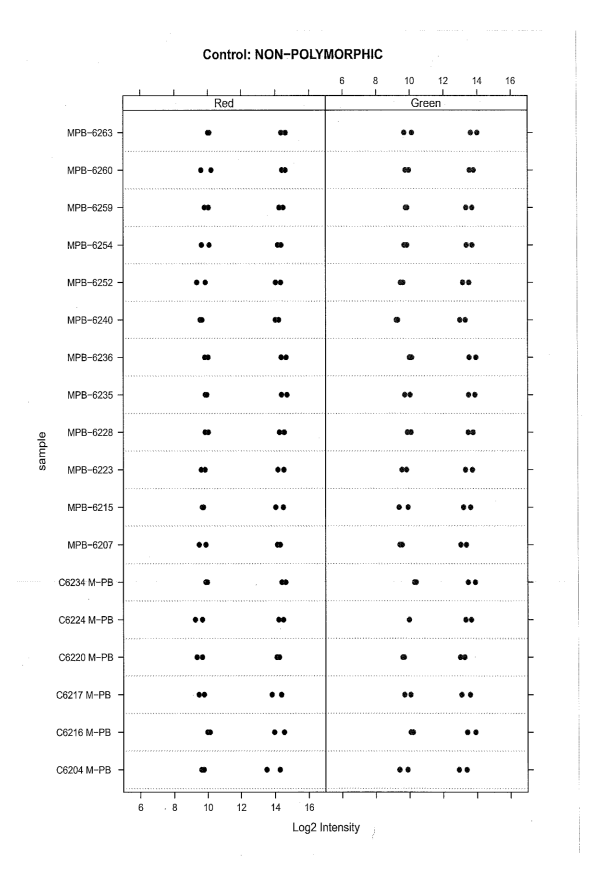
#### Control: BISULFITE CONVERSION I

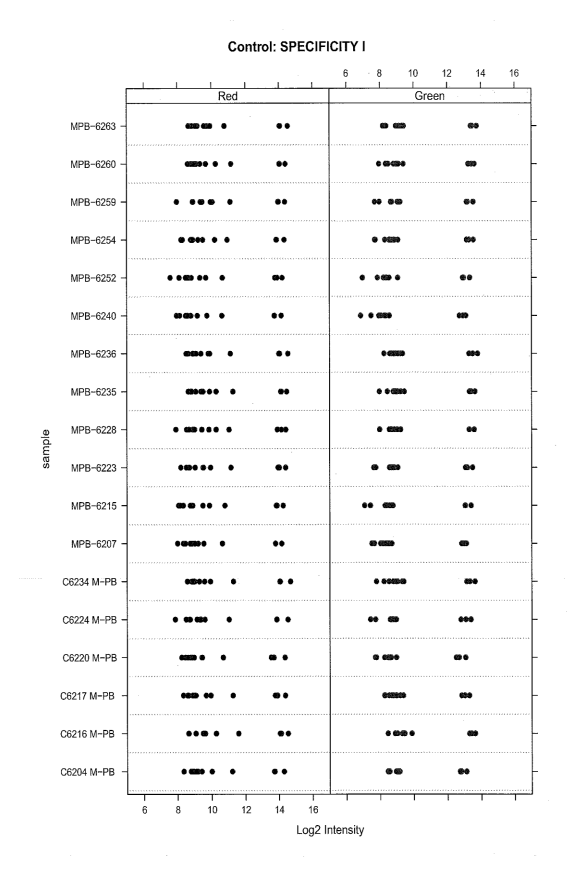


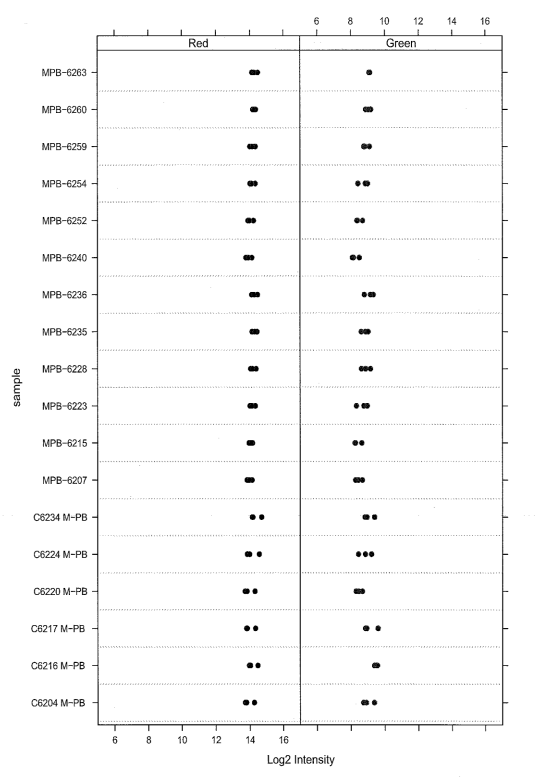
								6	8	10	12	14	16
		I	. I	R	ed		I			Gre	en		l
	MPB-6263 -				۲		8			63		•	•
	MPB-6260 -				٠		•		6	•		0	
	MPB-6259 -				•		•		6			6	
	MPB-6254 -			*****	•	*******	•		8			•	
	MPB-6252 -			, ; , , , , , , ,	•		•		0			•	
	MPB-6240 -				•		•		69			6	
	MPB-6236 -				•		•		(	••		•	•
	MPB-6235 -				.•		•		•	9		6	
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	MPB-6215 -				•		•	,.,.,.,	69			٠	, ,
	MPB-6207 -				•		•		69			•	
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	С6220 М-РВ -				•		•		•				
	C6217 M-PB -				•		•		•	9		٠	
	C6216 M-PB -				•		۲					•	
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		6	8	10	12	14	16 Log2 I		I		I	I	I



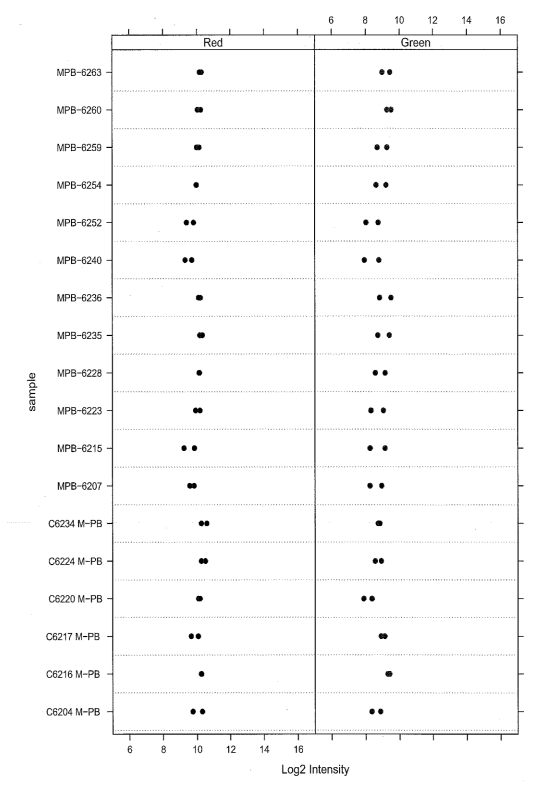
Control: HYBRIDIZATION







# Control: SPECIFICITY II



# Control: TARGET REMOVAL

Probe									
cg0446763		0.800366					cg0446763		
cg1116465			-0.46016			-	cg1116465		
cg0322139		0.73649	-0.4072				cg0322139		
cg0852247			-0.37087			U U	cg0852247		
cg1097852		0.806391	-0.34002	0.01313		-	cg1097852		
cg2413629		0.4966	-0.33572			-	cg2413629		
cg1563391		0.921117				-	cg1563391		
cg0897731		0.543341	-0.32363	0.006153		0	cg0897731		
cg0424670		0.682364	-0.31195	0.045569	0.999992	cg0424670	cg0424670	15809316	
cg2524308	0.149963	0.459685	-0.30972	0.000125	0.999992	cg2524308	cg2524308	55626484	
cg1255190	0.481229	0.786936	-0.30571	0.022987	0.999992	cg1255190	cg1255190	54759409	
cg0595694	0.534448	0.839984	-0.30554	0.00312	0.999992	cg0595694	cg0595694	57628353	
cg0851419	0.577935	0.876401	-0.29847	0.031578	0.999992	cg0851419	cg0851419	72705460	
cg2485165	0.346336	0.644359	-0.29802	0.031454	0.999992	cg2485165	cg2485165	22743423	
cg2138833	0.239326	0.531601	-0.29227	0.012766	0.999992	cg2138833	cg2138833	69638509	
cg2487392	0.488609	0.780509	-0.2919	0.000374	0.999992	cg2487392	cg2487392	29785495	
cg2616263	0.620694	0.910751	-0.29006	0.000618	0.999992	cg2616263	cg2616263	35653353	
cg0992568	0.461025	0.747655	-0.28663	0.027623	0.999992	cg0992568	cg0992568	36637470	
cg0120191	0.543413	0.827772	-0.28436	0.043415	0.999992	cg0120191	cg0120191	51627476	
cg1316715	0.659629	0.9436	-0.28397	0.021125	0.999992	cg1316715	cg1316715	52796390	
cg0112760	0.213465	0.497024	-0.28356	0.042571	0.999992	cg0112760	cg0112760	24715331	
cg2453864	0.356838	0.639609	-0.28277	0.00224	0.999992	cg2453864	cg2453864	56634317	
cg2742395	0.120616	0.402233	-0.28162	6.47E-06	0.999992	cg2742395	cg2742395	30689485	
cg0668491	0.501861	0.781294	-0.27943	0.038384	0.999992	cg0668491	cg0668491	70649349	
cg2129430	0.336143	0.61348	-0.27734	0.03464	0.999992	cg2129430	cg2129430	45781502	
cg0663177	0.333922	0.608484	-0.27456	0.048309	0.999992	cg0663177	cg0663177	24644310	
cg2453678	0.429272	0.700238	-0.27097	0.002489	0.999992	cg2453678	cg2453678	11672327	
cg2575542	0.162803	0.426214	-0.26341	0.031468	0.999992	cg2575542	cg2575542	14653467	
cg0559388	0.306954	0.56904	-0.26209	0.045719	0.999992	cg0559388	cg0559388	15800414	
cg1417593	0.363199	0.624853	-0.26165	0.011818	0.999992	cg1417593	cg1417593	47755467	
cg1880516	0.302811	0.564398	-0.26159	0.004211	0.999992	cg1880516	cg1880516	51795502	
cg1287021		0.723859	-0.26082	0.00802	0.999992	cg1287021	cg1287021	72752369	
cg0375488	0.445066	0.705665	-0.2606	0.022821	0.999992	cg0375488	cg0375488	45731391	
cg1783975	0.229936	0.490444	-0.26051	0.022773	0.999992	cg1783975	cg1783975	70682329	
cg2667988	0.236455	0.487702	-0.25125	0.048664	0.999992	- cg2667988	cg2667988	58642406	
cg2758679	0.429769	0.678816	-0.24905	0.035195	0.999992	cg2758679	cg2758679	12784395	
cg1536550	0.61212	0.859914	-0.24779	0.034901	0.999992	cg1536550	cg1536550	35642301	
cg1642309	0.454119	0.695547	-0.24143	0.042855	0.999992	cg1642309	cg1642309	42782411	
cg0777476			-0.24109	0.007408			cg0777476		
cg2497656		0.9144					cg2497656		
cg0450634							cg0450634		
cg1934487							cg1934487		
cg2394713							cg2394713		
cg1907951							cg1907951		
cg1337008						•	cg1337008		
cg0482944						-	cg0482944		
-90 1959-4	5.025001	5.550 105	0.22104	0.020011	5.55555E	-00,02044	-00.02044		

cg2369827	0.632711	0.842081			0.999992 cg2369827 cg2369827 70810465
cg1367916		0.787173	-0.20723	0.000851	0.999992 cg1367916 cg1367916 58762470
cg0448683		0.939273	-0.20504	0.04044	0.999992 cg0448683 cg0448683 61780346
cg2357685		0.709808	-0.20453	0.012287	0.999992 cg2357685 cg2357685 74615430
cg1541173	0.71671	0.921222	-0.20451	0.039182	0.999992 cg1541173 cg1541173 29723310
cg2693049		0.909556	-0.20434	0.003532	0.999992 cg2693049 cg2693049 36600316
cg1704224		0.581196	-0.20362	0.026914	0.999992 cg1704224 cg1704224 19652463
cg0382729		0.660819	-0.20096	0.01673	0.999992 cg0382729 cg0382729 74720388
cg1467135		0.375792	0.200115	0.027063	0.999992 cg1467135 cg1467135 33748352
cg0316740		0.648112	0.20056	0.02498	0.999992 cg0316740 cg0316740 41609406
cg2201856		0.753163	0.201689	0.043565	0.999992 cg2201856 cg2201856 14803393
cg2729222	0.826475	0.623284	0.203191	0.016989	0.999992 cg2729222 cg2729222 41607394
cg0457918	0.825164	0.62187	0.203293	0.046919	0.999992 cg0457918 cg0457918 11749430
cg1681031	0.65546	0.451768	0.203693	0.035013	0.999992 cg1681031 cg1681031 57768347
cg2420858	0.244437	0.040272	0.204165	0.002108	0.999992 cg2420858 cg2420858 52723466
cg1584536	0.27708	0.069441	0.207639	0.026749	0.999992 cg1584536 cg1584536 68737498
cg0146123		0.617264	0.207823	0.01631	0.999992 cg0146123 cg0146123 31654378
cg0979888		0.336846	0.210191	0.004471	0.999992 cg0979888 cg0979888 52724391
cg1233913		0.738073	0.210228	0.001594	0.999992 cg1233913 cg1233913 55756448
cg0230902		0.57016	0.213493	0.014514	0.999992 cg0230902 cg0230902 33758486
cg0027464		0.626229	0.2154	0.043587	0.999992 cg0027464 cg0027464 47758438
cg0191436		0.731168	0.217267	0.014629	0.999992 cg0191436 cg0191436 64666443
cg0768464		0.690243	0.217451	0.03079	0.999992 cg0768464 cg0768464 38691370
cg0865722		0.16906	0.217688	0.031716	0.999992 cg0865722 cg0865722 67723380
cg0352965		0.409944	0.218588	0.034733	0.999992 cg0352965 cg0352965 54618309
cg0255387		0.658326	0.220648	0.022832	0.999992 cg0255387 cg0255387 12768430
cg2676476			0.220806	0.010795	0.999992 cg2676476 cg2676476 37790457
cg1679183		0.63495	0.223285	0.02141	0.999992 cg1679183 cg1679183 37644405
cg1327992		0.602627	0.225377	0.015159	0.999992 cg1327992 cg1327992 49683476
cg0129687		0.340315	0.229877	0.024778	0.999992 cg0129687 cg0129687 29677416
cg0233894		0.725499	0.233625	0.026863	0.999992 cg0233894 cg0233894 24753391
cg0365105		0.574017	0.234234	0.024905	0.999992 cg0365105 cg0365105 21645475
cg0381217		0.548839	0.23494	0.046879	0.999992 cg0381217 cg0381217 53764329
cg1921470		0.269661	0.23506	0.047232	0.999992 cg1921470 cg1921470 51609466
cg1303102		0.427737	0.235071	0.013885	0.999992 cg1303102 cg1303102 61688421
cg0901915		0.21724	0.236393	0.026767	0.999992 cg0901915 cg0901915 73687380
cg2068168		0.57275	0.236752	0.006181	0.999992 cg2068168 cg2068168 21696495
cg1188103		0.468427	0.241189	0.034223	0.999992 cg1188103 cg1188103 24635354
cg0212689			0.24307		0.999992 cg0212689 cg0212689 21603365
cg0093943				0.003286	0.999992 cg0093943 cg0093943 70718348
0		0.670628			<b>u</b>
0		0.141706		0.002399	
cg0877964		0.557188			0
0		0.277221		0.011315	
cg1400175			0.258447	0.018637	0.999992 cg1400175 cg1400175 11758404
cg1301702			0.258672		
cg0938768	0.922428	0.663113	0.259315	0.03727	0.999992 cg0938768 cg0938768 47648483

 cg0014624	0.815754	0.554807	0.260947	0.0246	0.999992	cg0014624	cg0014624	44697466
cg2746787	0.693033	0.431724	0.261309	0.02545		cg2746787	•	
cg0289023	0.845479	0.581761	0.263718	0.049368	0.999992	cg0289023	cg0289023	64667483
cg1650756	0.936238	0.672067	0.264171	0.018053		cg1650756	Ŷ	
cg0999331	0.6496	0.372745	0.276855	0.039451		cg0999331	0	
cg0205655	0.723422	0.445196	0.278226	0.013002		cg0205655	0	
cg0600268	0.726262	0.447336	0.278926	0.023372	0.999992	cg0600268	cg0600268	24632477
cg1462190	0.745655	0.465995	0.27966	0.018646		cg1462190	•	
cg0183592	0.361672	0.078214	0.283457	0.001109	0.999992	cg0183592	cg0183592	21638313
cg0875045	0.599339	0.310648	0.288692	0.024058		cg0875045	Ģ	
cg0332735	0.74072	0.450433	0.290287	0.01305		cg0332735	•	
cg1193653	0.925907	0.634749	0.291158	0.033668	0.999992	cg1193653	cg1193653	24658402
cg1083223	0.855719	0.55978	0.295939	0.016736	0.999992	cg1083223	cg1083223	57802488
cg0724084	0.83221	0.53567	0.29654	0.025943	0.999992	cg0724084	cg0724084	35619467
cg1158502	0.713611	0.410317	0.303295	0.011149	0.999992	cg1158502	cg1158502	68619457
cg0810398	0.746795	0.442793	0.304002	0.020788	0.999992	cg0810398	cg0810398	29755444
cg2516514	0.775347	0.471297	0.304051	0.004235	0.999992	cg2516514	cg2516514	69721466
cg1220863	0.674749	0.366219	0.30853	0.041976	0.999992	cg1220863	cg1220863	27767440
cg2158700	0.748968	0.428555	0.320413	0.019786	0.999992	cg2158700	cg2158700	66612314
cg0580958	0.579108	0.258094	0.321014	0.019395	0.999992	cg0580958	cg0580958	47748455
cg2135833	0.758876	0.437735	0.321141	0.010715	0.999992	cg2135833	cg2135833	14702477
cg0044354	0.735279	0.402488	0.332791	0.012894	0.999992	cg0044354	cg0044354	12725423
cg0003321	0.541816	0.207825	0.33399	0.000666	0.999992	cg0003321	cg0003321	73611364
cg0461002	0.800062	0.46058	0.339482	0.013614	0.999992	cg0461002	cg0461002	16698312
cg1478255	0.92087	0.580606	0.340264	0.003323	0.999992	cg1478255	cg1478255	50621445
cg1122971	0.498728	0.154903	0.343825	0.011955	0.999992	cg1122971	cg1122971	42796477
cg0105569	0.728802	0.379156	0.349646	0.030341	0.999992	cg0105569	cg0105569	63796358
cg0509381	0.924233	0.574417	0.349816	0.004392	0.999992	cg0509381	cg0509381	68717455
cg0468351	0.66586	0.314219	0.35164	0.010373	0.999992	cg0468351	cg0468351	51686418
çg0978537	0.758268	0.367113	0.391156	0.007481	0.999992	cg0978537	cg0978537	41742374
cg2670559			0.4259	0.002307	0.999992	cg2670559	cg2670559	44686317
cg1234250	0.670831	0.242	0.428831	0.002112	0.999992	cg1234250	cg1234250	18752444
NA	NA	NA	NA	NA	NA	NA	NA	NA
NA	NA	NA	NA	NA	NA	NA	NA	NA
NA	NA	NA	NA	NA	NA	NA	NA	NA
NA	NA	NA	NA	NA	NA	NA	NA	NA
NA	NA	NA	NA	NA	NA	NA	NA	NA
NA	NA	NA	NA	NA	NA	NA	NA	NA

ΤΑΑΑCCTCCCTATTCTATTCCCTAAA ΙΙ			har Forward_Si Ger. דדדאאאקזינ	37	11	
CATAAACAAAATACACTCCTCTCAA <sup>-</sup> II			ATCCCGTT/	37	8	
TCTAATAAACATCTTCCCRAAATAACII			TGAAATTC	37	1	
TCTAATCACCTACAACCTTCTATCAA II			GCTCCCTT	37	3	
TTTATCCCTCCAACTACCCCAAACRC II			TTGACAAT.	37	7	
ACATTTCACAAAAACATACAAAAAT.II			TACCCTCA	37	11	
AAAACTAACRCRAAAACTTTCCTTTCII			ACAGCCAA	37	10	
			AATCTCTC/	37	3	
TTTACTCTAACACTAAACACAAAAT11			GTAAACAG	37	1	
TAACAAAT, 43698461 TAACAAAT, I	А	Red	TTTTCACA/	37	4	
	А	·	CTITITCAT	37	7	
			CCCAGGGA	37	8	
			GGTCTAGC	37	21	
			CCCTCAGC	37	11	
			TACAATGG	37	11	
			ACAGCTTG	37	11	
				37	7	
			CATGGTTT		19	
			TGGATCCA	37	19 15	
				37		
			ACAAGCTT	37	1	
		<b>D</b> 1	CCACCAGC	37	9	
ACACAAAA 29671432 ACACGAAAI	A	Red	GGGTCAG	37	8	
			AGGTCTGA	37	3	
ΤΑCΤΑΤΤΑΑCΤΑΑΑΑΑΑΑCΑΑCAAA ΙΙ			CTGTCACC,	37	19	
		<b>D</b> 1	GCTTAATT(	37	1	
AATATTTA/ 56680385 AATATTTA/1	A	Red	CCTGCCTC	37	12	
ACAAAAAA 35767456 ACGAAAAAI	A	Red	CTGTGCCC.	37	8 -	
CATTCAAA 70711339 CGTTCGAA I	C	Grn	GGCGGCTC	37	19	
ACTATATACTCCTTTCTACCTTCATA, II			CTGGGATA	37	7	
			GGAAACAG	37	14	
CAAAAAAC 49795427 CGAAAAACI	A	Red	TATATGAG	37	19	
TCCTAAAACAAATCAATTTCTAAAC/ II			GCCTGATA	37	14	
CTCCAATTTCTTTCTTTTATAACTAA(II			CCCACCAC	37	8	
ATTATTACCTCAACATCTTCCTTAAT, II			GCTCAAAA	37	21	
CAAATATACRATCATCAATAATCAC(II			CACTCGAC	37	10	
AAAACTTTCTTAACATAACCCTTAAAII			TGTACTGG	37	5	
TAATCAATAAACCTCCTTCATTACAC II			TTGTGCCA	37	6	
ΤΑΑΤΑΤΑΑΤΟΤΑΤΑΑΟΟΑΟΑΤΤΤΤΟ ΙΙ			TTCCATTAI	37	14	
ACCATAAA 21733390 ACCGTAAA	С	Grn	GCGATTTT	37	5	
TACACAAA 61624432 TACACAAA I	Т	Red	CTGATCTT	37	14	
ΤΤCCTTCRAATAAATACAAAAAACA/ ΙΙ			CTGGGATT	37	2	
ΤΤΤΟ ΑΑΤΤΟ ΑΑΤΑΑΑΤΑΑΤΙΙ			GATGACCA	37	12	
ΤΤCΑΑΑΑΤΑΑCΑΤΑCΑΑΤΑΑΤΑΑΤΑCΑ' ΙΙ			CGATGGCC	37	13	
TCTACTTCTCTATTCATATATATTTC/ II			GTTCTCAC <sup>2</sup>	37	17	
ACCCCATATAACCTACCTACATCAT#II			AAACAGG	37	1	
CRACTAAATTTTTCCTTTTTAATAA/II			TCAACACT	37	5	

 ACTCCTACCATCTCTTACRACTAATT II			CCTTTCCA1	37	10	 
AACTAAAT 60697445 AACTAAAT I	Т	Red	TCCGCAGC	37	6	
ΑΤΑΤΤΤΟΤΑΑΑΟΟΤΟΑΑΟΤΑΑΑΤΤΑΑΙΙ			GTTCCAGG	37	1	
CRATTAACAAAATACTAACAAAACA II			GAGGTTCT	37	5	
ΑΑΤCCCTCCTATAAAAACCAACTTC1			TGGCATTT,	37	12	
CACRTAAAATATAACCCACCCAAAA II			ACACCTCT	37	16	
ΤΟΤΤΑΑΑΑΑΑΑΑΤΑΟΟΑΟΑΤΑΟΤΑ΄ ΙΙ			ATGGCCAA	37	17	
AACRAAAAAACCTCAAAACTCRCCC II			AGGCCCGC	37 Y		
ΑCATAAATATTACAAACCTCTCTAA/ ΙΙ			TTGTATGA	37 Y		
ΑCAAATCTATAATACTTAAAACATC/ ΙΙ			GCTGTATG	-37	2	
ΤΤΑΑΑΑΑΑΑΑΑΑΤΑΤΑΤΑΑΤΑΤΑΑ			TGAGATGT	37	1	
ΤΑΤΑΑΤΟΟΟΑΑΑΑΑΤΟΟΟΤΑΑΑΟΟΑ			TCTTCAGA	37	6	
ΑΑCΑΑCΑCΤΑΑΑΑΑΑΤΤΑΑΤΑΑCTC ΙΙ			GGTTCTTT(	37	15	
AACCRAAATCAAAACCACATCCTCT II			GATAATCA	37	11	
ΑΑΑΑΑΑΤΤΑΤΟΤΤΤΤΑΑΑΑΑΑ			GAGGTCTC	37	7	
AATCTTCCCAATACAAAACTTAACA1II			CCAGCCAC	37	5	
ΑCCAACAATCAAAACTCCCTACTTC(			TTCTTTTCT	37	3	
AAAACCAA 33707303 AAAACCGAI	Т	Red	TGGGGGG	37	12	
ΑΑCΑΤΑΑΑCCACCRCACCTAACCTA			CTGTGCGA	37	15	
ΑΤΑΤΤΑΑΤ( 13769337 ΑΤΑΤΤΑΑΤ(Ι	С	Grn	GAAGGCA(	37	10	
AAACCCTTCRAAAAAACCRAAACAT II			GAGGAAA/	37	16	
ATATTTAATCTTCTCACACAAATTTT, II			GAGAAGA/	37	3	
CTACCAAACTACTATAATACRCCRA/ II			CAGGCTGC	37	16	
ΑΑΤΤΑΤΑΤΑΑΑΑΤΑΑΤΟΑΤΟΤΑΤΟΤΤΑΤ ΙΙ			CGTGGGCC	37	20	
TCCCTATCTTAATAAATCRATTCTAT II			CTATTGCA	37	3	
ΑΑΑCCTATTCCTATCTCTTTCTAAAA ΙΙ			TTGCAGCT	37	16	
CAAACCAC 47742450 CAAACCGCI	А	Red	GGCAGGG	37	16	
ΑΑΑΤCCΑΤΑCΑΑΑΑCCAAAACACCC ΙΙ			GTAGAATC	37	16	
ATTCAAAA 59679461 ATTCGAAA	А	Red	TAGGGCCC	37	7	
ΤΟΤΤΑΟΤΟΟΤΑΑΤΟΟΤΟΤΑΑΑΑΑΤΑΟ ΙΙ			CAGGCCCC	37	1	
CATTCRATTTTTTCCCTATATATTA/ II			CGGAACAA	37	2	
CRATATCTAAACAACRAAATATCTT(			CCCACTGT	37	13	
ΑCTCACTCAAAACTTTAATACATTTC ΙΙ			CAGGGAA(	37	7	
ΑΑΑΑCCAAAATCTTACTCAAAAATT, ΙΙ			TTTCTCCA(	37	7	
ΑΑΑΑΑΑΑΑΑΑΤΟΑΤΤΑΑΑΑΑΑΑΑΑΑΑΑ			GATGAGAG	37	7	
ΤCACACAACACCCTCTTAAAACAAA" ΙΙ			TTCCCACA	37	8	
ΤΤΑΤCCCTACAAACCCTAACTAAAC/			CGCTCACC	37	13	
<b>CRAATTAAATCCATTTTCAAACAAA( II</b>			TCCACATC(	37	6	
<b>CAAAAACAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA</b>			TGTAGAGC	37	11	
ΑΑΑCΑΑΑCΑΑCΑΑΑCΤΑCΤΑΤΑ ΙΙ			GTAGCCAG	37	14	
RAATCRTAAAAAATAAATTTCTCCA1II			GAGCGAC1	37 Y		
TCCCTTCTTATCTCTCTAATAACTTT/ II			ATGAGTTC	37	2	
ΤΑΑΑΑCTΑCTΑΑΑΤΑΑΑΑCCAAAAT ΙΙ			TTTCTCTCC	37	13	
TCCTTACCTACTAACTCCACCTACAT II			TCCGTGCT	37	6	
TCRCTTTTACTTATATATATAAATTAA II			ΠΤΑΠΟΠ	37	17	
ΑCCCAATCAATACCATTTATTATACA			GATGCAAA	37	5	
AACTTCCTCRCCTTTACCRTAACATT II			ACGCCAGC	37	10	

AATAC	CCACRATA	AACCACTAT	ITA'II			GGCACA	\G <i>F</i>	37	15	
TACTAC	TCCTCACT	AATACCTC	ACC II			GGCTGA	GC	37	8	
AAATCI	ACTITICTC/	АТААТСТАА	ITTC II			CCCCAG	GT	37	19	
CACTCT	AATAAAAT	CCACAAATA	ATA" II			GAGATO	6AC	37	6	
TAAAC	СТААААТС	RTATTCTAA	CRT II			TGGCCT	CC.	37	10	
AACATA	аааасатт	ATTCTCCCA	ACTIII			AATGAA	AG	37 Y		
AACTA	TC( 52784	353 GACTA	TTCII	А	Red	TGCGTG	AC	37	8	
AAAAC	CTTAAACCC	CCCATAACA	aaa II			GTATCG	GE	37	11	
TTCTAT	ACCAACRC	TTCAACTAA	CAA II			GTGAAG	GAC	37	11	
TTCCTA	CRTATITIC	TATCCAAA	TCC" II			TTGATT	CC1	37	17	
AATAA	AATAACCAA		CCT/II			TCTGGG	AA	37	1	
RACTA	сттстаатс	ACTAAATCA	ATTA II			TTGGTG	GA	37	11	
AAAAA	AAAACATA	ΑΑΤϹΤΑΑΑϹ	TAA II			GACTIT	ACI	37	14	
CAACA	АСТАТТАСТ	AATTCAACT	ITAA II			CCTTCTC	000	37	10	
AAATC	ATCTCCTT	CAACATCCT	°AAA II			CCACTT	CA(	37	5	
CTCCCT	сті 67753	457 CTCCCT	TCTTI	С	Grn	CGGAAA	ACC	37	17	
CCCCA		ATCCACTA	AAA II			ACTGAG	icg	37	12	
TTCAA	АССТАААА	AACRATAA	ATT/II			GAAAG	GC4	37	11	
AAAAA	атсаалаа	ACCTAATAC	CRT II			AAGACO	TA	37	10	
AAAAC	ТААААААС	АСАААССАА				TTCTGA	ΠC	37	21	
ATTCCT		ТАААААТСТ	FACC II			AGGCCC	CGC	37	17	
AAATTO	СТАААААСТ	AACCTAAA	ACT, II			AGCCTG	ΓAT	37	17	
AAAAA	TACRATAA/	ACRCCRAAA	TAA II			AGCCCT	GC	37	8	
TAAAA	CAACCCTAC	TCTAAAAA	CCA, II			CTTGCC	CCi	37	19	
AAAAT	CCRTCCTAC	TAAATTATC	CAA/ II			CTGGTG	στ	37	6	
CRATA	ΓΑΑCTACTT	ACAAAAAC	AAC, II			TGTATG	iGC	37	2	
CAATA	ACCTTATCT	AATTTATCA	ATTT II			TTCACT	TG1	37	3	
AAACA	TCAAAAAT	CAATTCCCC	AAT(II			ATACTA	AG	37	4	
AAAAA	AACTTAAA	TTTTACRTA	AAA(II			GAAACA	AA/	37	10	
ATAAT	ΑΤΑΑCΤΑΤΤ	CCACRACA	aac⁻ II			ACCATA	AT	37	15	
TAATC	RCTATCAAA	TATACRCCA	AAA(II			CTTGGC	CT	37	13	
CCAAT	AAAACAAC	AATCRCTAT	CAT II			GGGCA	TG/	37	2	
NA	NA	NA	NA	NA	NA	NA	NA	NA		
NA	NA	NA	NA	NA	NA	NA	NA	NA		
NA	NA	NA	NA	NA	NA	NA	NA	NA		
NA	NA	NA	NA	NA	NA	NA	NA	NA		
NA	NA	NA	NA	NA	NA	NA	NA	NA		
NA	NA	NA	NA	NA	NA	NA	NA	NA		

		~!			1.14.11.127
				and Probe_SNP Probe_SNP Randor	
14495049 CG			14451625 F	rs5566583 NA	NA
1.44E+08 AT			1.45E+08 R	NA	NA
2.48E+08 CG			2.46E+08 R	NA	NA
1.12E+08 CG			1.13E+08 R	NA	NA
1.43E+08 TT			1.43E+08 F	rs5827052: NA	NA
15180922 CA			15137498 R	NA	NA
3179703 CG			3169703 R	rs4881107 NA	NA
1.68E+08 CG		3		NA	NA
2.47E+08 CG			2.45E+08 F	rs3129547 NA	NA
40267141 TG			39943536 F	NA	NA
66785137 GG			66422572 F	rs9443 NA	NA
22133375 CG			22189320 F	rs6982089 NA	NA
46075092 CG			44899520 R	rs2838613 NA	NA
66362959 TT			66119535 R	, NA	NA
3606550 TC			3596410 R	rs3765725 NA	NA
14431708 CG			14388284 F	rs1102321; NA	NA
1.03E+08 CG			1.03E+08 R	TRU	
3373819 CG			3324819 F	rs1297412  NA	NA
28644585 CG			26318180 F	rs6200697' NA	NA
1562535 CG			1552398 R	NA	NA
1.29E+08 AC	CTTCCGG	-	1.28E+08 R	NA	NA
216453 AC			206453 R	rs5709037(NA	NA
1.27E+08 CG	GGGAGG		1.28E+08 F	NA	NA
1792217 GC	CTGTTGG		1743217 F	rs7258947 NA	NA
8120055 AA	AGTCTGA		8042642 F	rs7263424! NA	NA
1.33E+08 GA			1.31E+08 F	rs4076044 NA	NA
216659 GC	CGGAGG	8		NA	NA
13875111 CG	STTCGGG		13736111 R	rs3620361 NA	NA
77827379 CT			77665315 R	NA	NA
23018807 CG			22088647 R	NA	NA
36265700 CC	GAGAAAC		40957540 R	NA	NA
26862142 CG	GCTAAG		25931982 F	rs7359792 NA	NA
1.25E+08 CG	GGAAATT	8	1.25E+08 R	NA	NA
46077562 TT	ATTGCC	21	44901990 R	NA	NA
94136 CG		10		rs4607995 NA	NA
13664584 GC	GGCTTTC		13717584 R	rs870546 NA	NA
1.49E+08 GC	GTCAATG	6	1.49E+08 F	rs4897076 NA	NA
22279816 CG	GTGATCT	14	21349656 R	rs2856411 NA	NA
1.8E+08 AC	CCGTGAG	5	1.8E+08 F	rs1043480; NA	NA
24587638 CG	GGAAAC	14	23657478 R	rs7784824: NA	NA
1.6E+08 TC	CCTTCGG,	2	1.6E+08 F	rs1020453!NA	NA
1.31E+08 CG	GGCATGT	12	1.3E+08 R	NA	NA
1.15E+08 CG		13	1.14E+08 R	NA	NA
14936230 CT	IGCTTCT(	17	14876955 R	NA	NA
55260950 CC	CCCATGT	1	55033538 R	NA	NA
1.47E+08 G	GCTAAAT	5	1.47E+08 F	rs7700488 NA	NA

1.21E+08 CTCCTGCC/		1.21E+08				NA	
1.61E+08 CGGGAAG(	6	1.61E+08				NA	
1.61E+08 TATTTCTG(	1	1.6E+08	Ŕ			NA	
373299 GGTTGGCA		426299				NA	
9886905 CGGTGCTC		9778172		rs1084463.		NA	
71458897 ACGTGGAA	16	70016398	R	rs7629132:	NA	NA	
77136834 CTTGGGGA	17	74648429	R	rs2707040		NA	
24454927 GCGGAGG()	ſ	22864315	R		NA	NA	
7676811 CGTGAGTA Y	1	7736811	R		NA	NA	
2.41E+08 CAGGTCTG	2	2.41E+08	R	rs2352821	NA	NA	
1.12E+08 TGAGGGC4	1	1.12E+08	R	rs7737	NA	NA	
30980847 GTGATCCC	6	31088826	R	rs7527209	NA	NA	
88119834 CGGGCCAC	15	85920838	R		NA	NA	
66104993 GCCGAAGT	11	65861569	F	rs3116068	NA	NA	
22122872 GGGGACT(	7	22089397	R		NA	NA	
16785399 ATCTTCCC/	5	16838399	R		NA	NA	
1.91E+08 CCAGCAAT	3	1.93E+08	R	rs79001781	NA	NA	
1.08E+08 CGGGGCC/	12	1.07E+08	F		NA	NA	
90927939 CGATTGAA	15	88728943	F	rs9796504	NA	NA	
855060 CGGGGCCC	10	845060	R		NA	NA	
88238863 CGTTGGCT	16	86796364	F	rs1244663 <sup>-</sup>	NA	NA	
32511650 CGCTGTAT.	3	32486654	R	rs421653	NA	NA	
88296994 TGCCAGGC	16	86854495	R 💡	rs2873329(	NA	NA	
170641 GTTGTGTG	20	118641	R		NA	NA	
1.04E+08 CGCCCAAT.	3	1.06E+08	R	rs6226057!	NA	NA	
28273096 GGCCTATT	16	28180597	F	rs4788054	NA	NA	
87682142 CAGGCCGC	16	86239643	R		NA	NA	
73102243 AATCCATA	16	71659744	F	rs1183967	NA	NA	
1.57E+08 GTTCGGGA	7	1.57E+08	R		NA	NA	
2274955 CGAGGGGG	1	2264815	R		NA	NA	
2.25E+08 ATTCGGTT	2	2.25E+08	R	rs1686590	NA	NA	
50194643 GGTGTCTG	13	49092644	F		NA	NA	
44184403 CTCACTCA(	7	44150928	F	rs1330638	NA	NA	
3157722 CGCCTCCC	· 7	3124248	F		NA	NA	
1.08E+08 GAGGAAG(	7	1.07E+08	R		NA	NA	
19616280 CACACAGC	8	19660560	R		NA	NA	
1.13E+08 TGTCCCTG	13	1.12E+08	R		NA	NA	
1.54E+08 CGCATGGT	6	1.54E+08	R		NA	NA	
13938802 CGTGCATA	11	13895378	R	rs7131580	NA	NA	
55151579 GGCAGGC/	14	54221329	R		NA	NA	
14107195 CGTGCAGC	Y	12617195	F		NA	NA	
2.01E+08 CCCTTCTTA	2	2E+08	R		NA	NA	
50194554 CGCCTCTG	13	49092555	R		NA	NA	
1.57E+08 CCTTGCCT(	6	1.57E+08	R	rs7582559(	NA	NA	
64302651 CGGCCTAA	17	61733113	R		NA	NA	
1.4E+08 CCCAGTCA	5	1.4E+08	F	rs6849	NA	NA	
13688165 ACTTCCTC(	10	13728171	R		NA	NA	

62899159	GTGCTCCA	1	5	60686451	R			NA	NA
22266134	GCTGCTCC		8	22322079	R			NA	NA
55735946	GGTCTGCT	1	.9	60427758	F		rs1261080:	TRUE	NA
33158020	ACTCTGAT		6	33265998	F		rs7471695،	NA	NA
1.32E+08	GAGCTCTG	1	0	1.31E+08	R			NA	NA
8552374	CGGGAATC	Y_		8612374	F			NA	NA
80783997	CGGGGATI		8	80946552	R			NA	NA
65363274	GGGCCTTA	1	1	65119850	R			NA	NA
1.22E+08	TCTGTGCC	1	1	1.21E+08	F		rs1944694	NA	NA
6558815	TCCTGCGT	1	17	6499539	R			NA	NA
46979222	CGGGCCTC		1	46751809	R		rs4660355	NA	NA
1.34E+08	CGTCAAAG	1	11	1.33E+08	F		rs7303206	NA	NA
1.06E+08	GAAGGAG	1	14	1.05E+08	R			NA	NA
12438782	CGGGGAG	1	LO	12478788	R			NA	NA
1.1E+08	CGGGCAG1		5	1.1E+08	R			NA	NA
6558365	CGGAGCCA	1	17	6499089	R			NA	NA
670974	CGCCAAGC	1	L2	541235	F		rs3497962	NA	NA
66317822	TCAGGACC	1	11	66074398	R			NA	NA
839609	GAGAATCA	1	LO	829609	R			NA	NA
31709690	AGGCTGAG	2	21	30631561	R			NA	NA
6558440	CGCCTGAG	1	17	6499164	F			NA	NA
1645410	CGGGCTGC	1	17	1592160	F		rs6209005	NA	NA
1.44E+08	CGCACAGA		8	1.44E+08	R			NA	NA
8464538	GGGACAG(	1	19	8370538	R			NA	NA
33131893	CGGCTTTT		6	33239871	F		rs9405002	NA	NA
2.25E+08	GATGTAAC		2	2.25E+08	R			NA	NA
1.15E+08	CGGACTAT		3	1.17E+08	F		rs1093430	NA	NA
8728269 <b>7</b>	CGACAAAC		4	87501721	R			NA	NA
5047487	CGGCAGGT	-	10	5037487	R			NA	NA
60644157	CGGTGACC	-	15	58431449	F		rs1163365	NA	NA
1.12E+08	GGTCGCTG		13	1.11E+08	R			NA	NA
8530521	CGGAGGA		2	8447972	R			NA	NA
NA	NA	NA		NA	NA	NA	NA	NA	NA
NA	NA	NA		NA	NA	NA	NA	NA	NA
NA	NA	NA		NA	NA	NA	NA	NA	NA
NA	NA	NA		NA	NA	NA	NA	NA	NÁ
NA	NA	NA		NA	NA	NA	NA	NA	NA
NA	NA	NA		NA	NA	NA	NA	NA	NA

		UCSC_CpG_Relation_tcPhantom	DMR		HMM_Islar	
-	NM_01645 Body;Body			NA		
RHPN1	NM_05292 Body	chr8:14445 S_Shore		NA	8:1445288(	
		chr1:2478C S_Shore		NA	1:2458702(	
	NM_00100 Body;Body	;Body		TRUE		
	NM_17743 TSS200			NA		
INSC;INSC	NM_00103 Body;Body			TRUE		
		chr10:318C N_Shore		NA	10:316969:	
C3orf50	NR_02148! Body			TRUE		
CNST;CNST	NM_00113 Body;Body	chr1:24678S_Shore		TRUE		
				TRUE		
	NM_02290 3'UTR			NA		
-	NM_01806 5'UTR;1stE			NA		
KRTAP12-4	NM_19869 TSS1500;B			NA		
CCS	NM_00512 Body	chr11:663€ S_Shelf		NA		
TP73;TP73	; NM_00112 TSS1500;T	chr1:3607CN_Shore		NA ·		
				TRUE		
	I NM_17305 Body;Body			NA		
NFIC;NFIC	NM_00559 Body;Body			NA		
		chr15:2864 N_Shelf		NA		
MIB2;MIB2	NM_00117 Body;Body	chr1:1563€ N_Shore		NA	1:1551913-	
LMX1B	NM_00231 Body	chr9:12937 Island	DMR	TRUE	9:1284157	
		chr8:21635 Island		NA	8:206391-2	
				TRUE		
ATP8B3	NM_13881 Body	chr19:1795 N_Shelf		NA		
				TRUE		
		chr12:1329 Island		NA	12:131482:	
		chr8:21635 Island		NA	8:206391-2	
MRI1;MRI	1 NM_00103 TSS1500;T	Schr19:1387 Island		NA	19:137360!	
MAGI2	NM_01230 Body			TRUE		
				TRUE		
SNX26	NM_05294 TSS1500	chr19:362€ N_Shore		NA		
				TRUE		
				TRUE		
C21orf29;I	K NM_14499 Body;TSS1	500		NA		
TUBB8;TU	E NM_00116 Body;Body	chr10:9452 N_Shore		NA	10:83977-8	
				TRUE		
UST	NM_00571 Body			TRUE		
				NA		
BTNL9	NM_15254 3'UTR	chr5:18048 S_Shore		NA	5:1804187(	
DCAF11;D	( NM_18135 Body;Body	/ chr14:2458 S_Shelf		NA		
BAZ2B	NM_01345 5'UTR			TRUE		
		chr12:1313 N_Shelf		NA		
RASA3	NM_00736 Body	chr13:1147 Island		NA	13:113800	
				TRUE		
ттс22;πс	X NM_00111 Body;Body	1		TRUE		
SPINK5;SP	I NM_00112 Body;Body	/;Body		NA		

TIAL1;TIAL	NM_00103 Body;Body	1			TRUE		
					NA	6:1611784:	
C1orf192	_				NA		
AHRR	NM_02073 Body	chr5:37384 N_Shore			TRUE		
CLECL1	NM_17200 TSS1500				NA		
		chr16:7145 N_Shore			NA		
HRNBP3	NM_00108 5'UTR	chr17:7713S_Shelf			NA		
	ENM_15258 TSS200;TS	SchrY:24454 Island			NA	Y:22864174	
TTTY12	NR_00155: Body				NA		
		chr2:2412€ N_Shore			NA		
ADORA3;A	NM_00108 3'UTR;3'U	TR			NA		
					NA		
	) NR_02686! TSS1500				NA		
BRMS1;BR	INM_015393'UTR;3'U	T chr11:661C S_Shelf			NA		
		chr7:22122 Island			TRUE	7:2208893	
MYO10	NM_01233 Body				TRUE		
CCDC50;CO	CNM_17833 Body;Body	/ chr3:19104S_Shelf			NA		
		chr12:1082S_Shore			NA	12:106763	
		chr15:9093 N_Shelf			NA		
		chr10:855€ N_Shore			NA		
		chr16:8823S_Shore			NA	16:867963!	
		chr3:32509 S_Shelf			NA		
		chr16:8829S_Shelf			NA	16:868542:	
DEFB128	NM_00103 TSS1500				NA		
					TRUE		
		chr16:2827S_Shelf			TRUE		
JPH3	NM_02065 Body	chr16:8767S_Shelf			NA	16:862395(	
		chr16:7309 S_Shore			TRUE		
PTPRN2;P	T NM_00284 Body;Bod	y;Body			NA	7:1571566	
MORN1	NM_02484 Body	chr1:22763 N_Shore			NA		
FAM124B;	INM_02478 3'UTR;3'U	TR			NA		
					NA	13:490924:	
GCK;GCK;C	G NM_03350 3'UTR;3'U	T chr7:44184 N_Shore			NA		
					TRUE		
LAMB1	NM_00229 Body				TRUE		
		chr8:19614 S_Shore	C	DMR	NA		
		chr13:1128 Island			NA	13:111896!	
OPRM1;0	P NM_00114 Body;Bod	y;Body;1stExon;Body;Bo	ody;5'UTR;Bod	y;Body;E	Bi NA		
					TRUE		
SAMD4A;S	S, NM_00116 Body;Bod	y			TRUE		
		chrY:14107 Island			NA	Y:1261713	
C2orf69	NM_15368 Body	chr2:20077S_Shelf			NA		
					NA	13:490924:	
ARID1B;AF	R NM_01751 Body;Bod	y;Body			TRUE		
PRKCA	NM_00273 Body	chr17:6429S_Shelf			NA		
ZMAT2	NM_14472 3'UTR				NA		
FRMD4A	NM_01802 3'UTR		low-CpG:137	28022-1	3 NA		

 							TRUE		
SI C20A1/		35 Body;Bod	w Body Bo	du			NA	8:2232190	
SEC35A14	F, INIVI_013	55 BOUY,BOU	у, БОЦУ, БС	Juy			NA	19:604274(	
COI 1142	·C NM 080	67 Body Bod	v chr6·33	159 N Shore			NA		
MGMT	NM 002		1. 0111 0150				TRUE		
TTTY18	-	5(TSS1500					NA		
							NA		
KCNK7;KC	CNNM 033	34 1stExon;5	'l chr11:6	536 N Shelf			NA		
	-						TRUE		
			chr17:6	559 N_Shore			NA	17:649869	
DMBX1;D	NM_147	19 3'UTR;3'U	TR				NA		
			chr11:1	337 N_Shelf		RDMR	NA		
			chr14:1	061 N_Shore			NA		
CAMK1D;	C NM_020	39 Body;Bod	у				TRUE		
WDR36	NM_139	28 TSS1500	chr5:11	042 N_Shore		CDMR	NA		
			chr17:6	558 Island			NA	17:649869	
B4GALNT	3 NM_173						NA	12:541125-	
ACTN3	NM_001	10 Body		631S_Shelf			NA		
			chr10:8	39£ Island			NA		
KRTAP27	-1 NM_001	07 1stExon					NA		
				558 Island			NA	17:649869	
	· -	93 TSS1500;"					NA		
	NM_052	•		439 Island			NA		
RAB11B	-			464 N_Shore			NA		
	-	67 Body;Bod	y cnr6:33	1252_Shelf			NA TRUE		
CUL3	NM_003	,	TI chr2.11	537 N_Shore			NA		
	_	98 TSS1500;					NA		
-	_	13 TSS1500;					NA		
		13 Body;Bod					TRUE		
ANAA2,A	N. NIVI_001	.13 D007,500		117 Island			NA	13:110543!	
			01112.1	117 (Sland			TRUE	10.1100 10.	
NA	NA	NA	NA	NA	NA	NA	NA	NA	
NA	NA	NA	NA	NA	NA	NA	NA	NA	
NA	NA	NA	NA	NA	NA	NA	NA	NA	
NA	NA	NA	NA	NA	NA	NA	NA	NA	
NA	NA	NA	NA	NA	NA	NA	NA	NA	
NA	NA	NA	NA	NA	NA	NA	NA	NA	

Regulatory Regulatory DHS NA )3-144528896 NA 50-245870330 NA NA NA 11:151807 Unclassifie NA TRUE 1-3169822 NA NA 4:4026692 Unclassifie NA 1552467 NA 32-128417915 NA :06774 NA NA NA NA 196-131483014 NA 206774 NA 53-13736951 NA NA NA 19:362656 Unclassifie NA NA NA 21:460773 Unclassifie NA 4317 NA TRUE NA NA 51-180419744 NA NA NA NA 13:114782 Gene\_Asso TRUE NA TRUE NA

	NA
11-161178494	NA
	NA
	NA
	NA
16:714587 Unclassifi	
4 33864503	NA
1-22864502	NA NA
	NA
7:2212206 Promoter	
,,	NA
	NA
202-106763970	NA
	NA
	NA
56-86796487	TRUE
	TRUE
17-86854496	NA
	NA
	NA
	NA
05-86239722	TRUE
	TRUE
7:1574641 <sup>,</sup> Unclassifi	
	NA
	NA
10-49092680	NA
	NA
	NA
8:1961507: Promoter	NA r NA
577-111897215	NA NA
3//-11103/213	NA
	NA
	NA
3-12617480	NA
1 12017 100	NA
10-49092680	NA
	NA
	NA
	NA
	NA

3-2	2322173	NA	
	-60427866	NA	
		NA	
11	L:653626 Unclass	ifie TRUE	
		NA	
17	7:655871: Unclass	ifie NA	
1:	4697902 Unclass	ifie NA	
		NA	
		TRUE	
		NA	
5:	1104272 Promot	er_ NA	
17	7:655834 Unclass	ifie NA	
-54	1330	NA	
		NA	
		NA	
		NA	
17	7:655834 Unclass		
		NA	
		TRUE	
		NA	
55	4-110543657	NA	
		NA	
N		NA	
N		NA	
N.		NA	
N. N		NA NA	
		NA	
N.	a na	NA	

Appendix H Preliminary Data Differential DNA Methylation in Genes Associated with Immune Function in <sup>1</sup>University of North Dakota, Grand Forks, ND; <sup>2</sup>The Ohio State University, Columbus, OH Pregnant Women with Group B Streptococcus Colonization Michelle L. Wright<sup>1</sup>, Cindy M. Anderson<sup>2</sup>, and Joyce E. Ohm<sup>1</sup>



## **Introduction**

- Group B streptococcus (GBS) is the leading cause of infectious neonatal morbidity and mortality in the United States
- Maternal GBS colonization is the primary risk factor associated with neonatal infection
  - Ambiguous maternal risk factors for colonization
- 10-30% of pregnant women are colonized with GBS
- can be altered by environmental exposures resulting DNA methylation is an epigenomic modification that
  - Altered gene expression <u>\_</u>
    - Predisposition to disease
      - Purpose:
- Determine feasibility of evaluating DNA methylation patterns as a contributing factor to GBS colonization

### Methods

- DNA methylation in maternal peripheral white blood cells collected in first trimester of pregnancy were analyzed using Illumina Infinium®
  - Group differences in DNA methylation patterns in women with and without GBS colonization were
- CpG Dinucleotides change in gain or loss in methylation, respectively. Delta-beta criteria >0.2 or <-0.2 to determine</li> determined with:
  - Significance determined by t-test of mean methylation difference, p<0.05
    - Functional relevance of differentially methylated genes was determined by:
- GeneCards®, DAVID Bioinformatics Resources v6.7 (Huang, et al., 2009), PubMed
  - Utilized "high" classification stringency
- Lempicki, R. a. (2009). Systematic and integrative bioinformatics resources. Nature protocols, 4(1), 44–57. doi:10.1038/nprot.2008.211 Reference: Huang, D. W., Sherman, B. T., & analysis of large gene lists using DAVID

## **Preliminary Results**

- Results are from a secondary analysis of DNA methylation were used to determine feasibility for a larger study
- Preliminary results of differential DNA methylation resulted in >1,000 >20% difference in methylation in GBS positive (n=2) versus GBS CpG (phosphate linked cytosine-guanine base pairs) sites with
  - 350 CpG sites had >30% difference in methylation (Figure 1) negative (n=4) women
- Differential methylation most strongly associated with immune function pathways (Table 1)

# Figure 1. Potential DNA methylation biomarkers associated

dinucleotides with potential differential methylation in maternal peripheral blood delta beta > 0.3 or -0.3 are shown for 4 GBS negative maternal DNA CpG (phosphate linked cytosine-guanine base pairs) with maternal GBS status. Average beta scores from 350 and 2 GBS positive women GBS -

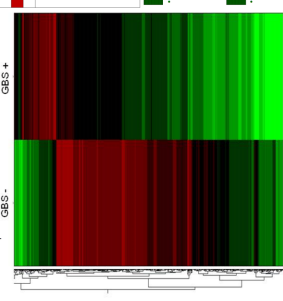


Table 1.	Functional an	Table 1. Functional analysis of differential DNA methylation
Cluster	Enrichment Score	Gene members in Functional cluster
		Methylation Loss
		MHC protein complex
		Antigen processing and presenting
-	3.96	Immunoglobulin C1-set
		Immunoglobulin/MHC, conserved site
		Immune response
		MHC class II receptor activity
		MHC class II, alpha/beta chain, N-terminal
ç	01 0	MHC class II protein complex
N	0.12	Antigen processing and presenting
		Intestinal immune network for IgA production
		Cell adhesion molecules
		Methylation Gain
٢	2.41	Pleckstrin homology
		MHC class II receptor activity
		MHC class II, alpha/beta chain, N-terminal
		MHC protein complex
		Immunoglobulin C1-set
2	2.31	MHC class II protein complex
		Antigen processing and presenting
		Cell adhesion molecules
		Intestinal immune network for IgA production
		Immune response
	Ŭ	Conclusions
Prelim	Preliminary data suggests:	ests:
• GBS	erentially methyli S colonization ar	Differentially methylated genes between women with and without GBS colonization are tightly clustered in immune function
path	pathways	

- immune function may contribute to GBS colonization susceptibility Differentially methylated CpG dinucleotides associated with

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