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UNIVERSITY OF ALBERTA

Layers of Epigenetic Regulation in the Transposed Imprinted Genes in the Prader-Willi Syndrome Domain

by

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A thesis submitted to the Faculty of Graduate Studies and Research in partial fulfillment of the requirements for the degree of Doctor of Philosophy in

Medical Sciences - Medical Genetics

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ABSTRACT

Not all genes are inherited equal. For parts of the mammalian genome, the two alleles inherited from the parents are made non-equivalent by epigenetic means. One allele is transcribed preferentially over the other, which is based on the gender of the parent contributing that allele. Human chromosome 15q11-q13 and mouse chromosome 7C contain large, syntenic imprinted domains. Previous work has established a model by which some of these genes are imprinted in a transcription dependant manner initiating at the imprinting center (IC). The transposed imprinted genes, *NDN/Ndn*, *MAGEL2/Magel2*, *MKRN3/Mkrn3* and *Frat3* are also paternally expressed and under control of this IC, although the mechanisms of this control have been less characterized.

To study imprinting of the transposed imprinted genes, a strategy was employed to examine tissue and allele-specific regulation at different scales. *NDN/Ndn* serves as an excellent representative for this cluster of co-regulated genes. Comparative analysis of the promoter sequences across different species gave clues as to the motifs involved in regulation of *NDN/Ndn*. Detailed examination of human *NDN* indicated tissue and allele-specific differences in accessibility of the promoter to *trans*-acting factors. The basis of this difference may lie in the differential chromatin context as evidenced by differences in histone modifications. Using chromatin immunoprecipitation (ChIP), I found that certain modifications correlated with expression of *NDN*, and that other marks are associated with allelic identity. In the search for a mechanism that potentially leads to allelic differences in chromatin modifications that exist regardless of tissue type, the involvement of a protein already shown to be important in imprinting was ascertained. This protein, CTCF, binds at

positions flanking the transposed imprinted genes in the mouse, and more interestingly, does so in an allele-specific manner.

These studies provide important clues about the layers of regulatory mechanisms in the tissue and allele-specific regulation of *NDN/Ndn* and the other transposed imprinted genes. From this work, I propose a model of regulation for the transposed imprinted genes that involves CTCF associated chromatin changes that lead to differential histone modifications and ultimately, to accessibility and function of individual promoters to bring about correct spatiotemporal and imprinted expression.

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LIST OF ABBREVIATIONS, SYMBOLS AND NOMENCLATURE

AS – Angelman Syndrome

BWS - Beckwith Wiedemann Syndrome

- ChIP chromatin immunoprecipitation
- DMD differentially methylated domain
- DMR differentially methylated region

EMSA - electrophoretic mobility shift assay

FB - fibroblast cell line

FISH - fluorescence in situ hybridization

IC - imprinting center

ICR – imprint control region

HAT – histone acetyltransferase

HDAC - histone deacetylase

HMT – histone methyltransferase

LCL – lymphoblast cell line

LCR - locus control region

MBD – methyl binding domain

PBS – phosphate buffered saline

PWS – Prader-Willi Syndrome

RNAi-RNA interference

SRO - smallest region of overlap

TF-transcription factor

UPD – uniparental disomy

ZF - zinc finger

Chapter 1
 State of the Art

Genomic Imprinting

It is fortunate that Gregor Mendel chose the humble pea plant for his Treatises on Plant Hybrids of 1865. The observations with which he discovered his three laws of heredity form the basis of genetics as we know it today. However, had his focus been on a mammalian model, imprinted inheritance may have wreaked havoc on his theories.

Genomic imprinting is a phenomenon by which alleles of certain genes are regulated differentially depending on the parental gender of origin (da Rocha and Ferguson-Smith 2004). For most genes in a mammalian genome, parental contributions of each allele are not overtly identified with respect to the parental gender from which it originated. Both alleles are free to exert their influence according to Mendelian laws. Imprinted genes, on the other hand, are inherited genetically from both parents, but each carries an epigenetic memory of the gender of the previous generation. This allelic identity causes the silencing of one allele in the offspring and breaks Mendel's first law by causing a functional hemizygous state for imprinted loci (independent dominance and recessiveness of alleles are made irrelevant).

For example, a trait that depends on an imprinted locus being expressed from the paternal allele and silent on the maternal allele (maternally imprinted) can appear to "skip" generations indefinitely through the maternal side of a family and reappear only when passed through a male germline. Thereafter, it can manifest itself if passed through the paternal side. This inactivation and activation of imprinted genes is done without changes in DNA sequence. Instead, allelic identity is kept epigenetically, that is, by modifications carried on or in association with DNA itself. These epigenetic marks are erased and written

differentially in oocyte and sperm precursors, giving rise to differential allelic identity in the offspring.

As far back as 1965, inheritance and effects of mutation in an imprinted gene, Igf2rwere characterized (Johnson 1974). The gene responsible had not been identified and was called the T locus, and instead of imprinting, the observations were considered within the nebulous phenomenon given the label of maternal effects (Haig 2004). Subsequently, debate raged as to the mechanistic defect in the T locus mutation; was it a cytoplasmic defect of the egg or maternal pronuclear defect? This mirrored a similar debate prior to 1984 regarding why parthenogenotes (embryos with uniparental genetic contribution) fail in development; was it a cytoplasmic deficiency in contribution from the sperm, or a nuclear genetic defect of the genome wide homozygous state? Both debates were settled in back-to-back papers in the April 5th issue of Nature, showing non-equivalence of parental genomes (Surani et al. 1984). The nuclear defect of the T locus lethality was also attributed to differential modification of parental genomes, but more specifically, of the locus on Chromosome 17 (McGrath and Solter 1984b). The two groups delineated imprinting as a field of study with seminal work on parthenogenetic and androgenetic mouse embryos (McGrath and Solter 1984a; Surani et al. 1984). Surani et al. used activated haploid eggs and added pronuclei from fertilized eggs to produce gynogenotes, while McGrath and Solter transplanted pronuclei between fertilized eggs to produce gynogenotes and androgenotes. These invariably failed to develop to term and showed the requirement of paternal and maternal contributions for normal development. While "epigenetics" was not a term coined for its current use back then, these studies implied the most fundamental

concepts of imprinted epigenetic memory of the previous generation, and the switch of that memory through the germline without genetic changes.

In humans, similar phenomena occur spontaneously in the form of hydatidiform moles and ovarian teratomas, containing only paternal and maternal genomes, respectively (Mowery-Rushton et al. 1996; da Rocha and Ferguson-Smith 2004). Hydatidiform moles consist of extraembryonic-like tissues while ovarian teratomas resemble tissues from the three germ layers. Histologically, these human uniparental tissues give telltale signs of a possible reason for the evolution of genomic imprinting, the sexual conflict model (Haig and Graham 1991). This model posits that the evolutionary driving force for fixation of imprinting is due to the unique maternal-offspring relationship in mammals. The disparity between maternal and paternal resources spent on a placental fetus pressures the paternally derived genome to increase growth of the fetus at the cost of future fecundity of the female, while the maternal genome has an interest in keeping growth in check for the sake of future fecundity. Consistent with this, hydatidiform moles consisting of paternal contributions are biased towards extraembryonic tissue growth, which would later increase nutrient transfer from mother to fetus.

Also of great interest is the mechanism by which imprinting arose. Similarities to X-inactivation are undeniable, and parallels have given great insight into the functions of both processes. X-inactivation is the process by which one X-chromosome in female cells is inactivated to achieve dosage compensation with respect to the male complement of a single X (Lyon 1961). While random X-inactivation and relevance to disease has received the most attention, it may be the exception and not the rule as X-inactivation is imprinted in

marsupials as well as extraembryonic tissues of some mammals. It has been postulated that the ancestral form of X-inactivation is imprinted, and that autosomes gained this characteristic through translocations from the X, and gave rise to autosomal imprinting (Lee 2003). Perhaps ancestral forms of one of the following imprinted genes or clusters will be found on the X-chromosome of a mammalian/marsupial ancestor. Debates continue regarding the role of imprinting in evolution, but it is clear numerous loci in the mammalian genome exhibit this phenomenon (Morison and Reeve 1998).

H19/Igf2

No comprehensive discussion of imprinting can omit mention of the *H19/IGF2* imprinted genes. They have become the prototype of all imprinted genes and is the most thoroughly characterized (Arney 2003; Delaval and Feil 2004). Located on mouse chromosome 7 and human chromosome 11, *H19* and *IGF2/Igf2* are reciprocally imprinted, being expressed from the maternal and paternal alleles, respectively (Figure 1-1 A) (Bartolomei et al. 1991; DeChiara et al. 1991). Imprinting of these two genes is controlled by an imprint control region (ICR) located between the two genes. Deletion of this *cis*acting element results in loss of imprinting and biallelic expression of both genes. Consistent with the conflict theory of imprinting discussed above, *Igf2* is a growth factor, and biallelic expression results in an increase in growth (Thorvaldsen et al. 1998), consistent with previous knock-out studies (DeChiara et al. 1991). The function of the ICR depends on a differentially methylated region (DMR, or sometimes DMD for differentially methylated domain), which is methylated on the paternal allele and unmethylated on the maternal allele. A methylation-regulated boundary model has been postulated whereby

methylation of the DMR upstream of *H19* allows enhancers to activate *Igf2* on the paternal allele and hypomethylation of the DMR activates a boundary element precluding enhancers from activating *Igf2*, which are now accessible to *H19* (Figure 1-1 A) (Schmidt et al. 1999; Delaval and Feil 2004). There are other allelic differences thought to contribute to allelic expression. Histone modifications for example have been found to be differential, with the unmethylated maternal allele of the ICR enriched for active modifications such as acetylation (Hu et al. 1998) and histone methylation at lysine 4 of H3, while the paternal allele carried histone methylation at lysine 9 of H3, a modification indicative of silent chromatin (Yang et al. 2003). Nuclease sensitivity, a measure of chromatin conformation also shows a more accessible maternal allele at the DMR (Hark and Tilghman 1998). Replication timing has also been found to be differential between the two alleles (Bickmore and Carothers 1995).



Figure 1-1. Mechanism of imprinting at Igf2/H19.

(A) Schematic of the methylation-regulated boundary model. Igf2 out-competes H19 for function of a distal enhancer on the paternal allele. CTCF is able to bind the unmethylated ICR of the maternal allele and block the enhancer from Igf2, allowing it to function at H19. (B) Intralocus loops in Igf2/H19 imprinting. The methylated paternal ICR associates with DMR2 of Igf2 to facilitate its expression. The CTCF bound maternal ICR associates with DMR1 of Igf2 and shifts it to a silent nuclear domain.

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Much has been learned since about the mechanisms of *H19/Igf2* imprinting, and this model has been elaborated in recent years. Most significantly, it has been found that CTCF, a protein that participates at almost all characterized mammalian insulators (Bell et al. 1999), functions in *H19/Igf2* imprinting (Bell and Felsenfeld 2000; Hark et al. 2000). Within the hypersensitive sites of the DMR (Hark and Tilghman 1998), are 4 CTCF binding sequences. These were found to have insulating activity in reporter assays, and binding was abrogated by DNA methylation. By binding the unmethylated maternal allele, CTCF acts to insulate *Igf2* from enhancers downstream of *H19*, thereby fulfilling the enhancer competition model mechanistically (Figure 1-1 A).

Furthermore, the mechanism of locus organization has also been elucidated. In elegant experiments involving two complementary lines of evidence, the higher order structure of the imprinted *H19/Igf2* genes have been characterized (Murrell et al. 2004). The data showing intralocus association imply that allele-specific long range loops are formed between the *H19* DMR and two other DMRs located at the 5' and 3' end of *Igf2* (DMR1 and DMR2, respectively). The model is that the *H19* DMR associates with DMR1 and DMR2 on the maternal and paternal alleles respectively, forming allele-specific epigenetic switch that shifts *Igf2* in and out of silent chromatin domain (Figure 1-1 B). How this aids in the access of enhancers in separated loop domains remains unclear and satisfactory convergence with the methylation-regulated boundary model is still lacking (Kato and Sasaki 2005).

BWS and the Kcnq1 locus

Adjacent to the H19/Igf2 imprinted genes, lies another imprinted locus. Beckwith-Wiedemann Syndrome (BWS) results from paternal uniparental disomy (UPD) of chromosome 11p15, disrupting both imprinted regions (Weksberg et al. 2003). Genes in both regions are candidates for phenotypes in BWS; therefore, to avoid confusion, this second imprinted locus will herein be referred to as the Kcnq1 locus. Five genes are imprinted in this locus: TSSC3, SLC22A1L, CDKN1C, KCNQ1 and LIT1 (seven in mouse with addition of Mash2 and Tssc4). With the exception of LIT1, all are maternally expressed. The ICR for this cluster is the KvDMR, located in an intron of KCNQ1, and this ICR controls expression of LIT1, which is an antisense transcript to KCNQ1 (Smilinich et al. 1999; Horike et al. 2000; Fitzpatrick et al. 2002). Imprinting of the Kcnq1 locus is independent of the H19/Igf2 locus (Caspary et al. 1998; Fitzpatrick et al. 2002). The mechanism of imprinting also appears to be quite different than this neighboring locus. Expression patterns do not support an enhancer competition model (Mancini-DiNardo et al. 2003). It has been shown however, that the unmethylated KvDMR functions as LITI's promoter and down-regulates other imprinted genes in cis on the paternal allele. The antisense nature of *LIT1* and the observation that *LIT1* is the only transcript expressed only on the paternal allele is reminiscent of the relationship between XIST and X-inactivation (Delaval and Feil 2004). In this system, XIST is transcribed from the silent X, and physically coats and silences that X in cis (Shibata and Lee 2004). Whether a similar mechanism exists for LIT1 is unclear, however it is tempting to speculate that such a mechanism exits. Even within these neighboring examples of imprinted domains on the

same chromosome, it is clear major differences exist in the mechanism of allelic regulation. Other imprinted domains such as *Igf2r*, *Callipyge* (not discussed here, but for review, see (Haig 2004)) and the PWS cluster, share this common theme of uncommon mechanisms.

PWS cluster at 15q11-13

A large cluster of imprinted genes resides on human chromosome 15q11-q13, with a region of conserved synteny on mouse chromosome 7C. In human, disruption of this region results in Prader-Willi syndrome (PWS) or Angelman syndrome (AS) (Nicholls and Knepper 2001). The determining factor between the two syndromes is parental origin of the remaining chromosome or region (Knoll et al. 1989). PWS results from a lack of paternal genetic contribution of the region either through maternal UPD (20%-30% of cases) or paternal deletion (~4Mb, 65%-75% of cases). In up to 5% of cases, an imprinting mutation causes assignment of a maternal identity to the paternal allele in the male germline. Angelman results from lack of maternal contribution by reciprocal chromosomal mechanisms as PWS, or by mutations in UBE3A (10%), a maternally expressed gene responsible for most if not all the features of AS (Burger et al. 2002). No single gene responsible for the neurobehavioral features of PWS, such as hypotonia, developmental delay and hyperphagia (Holm et al. 1993), has been identified (Lee and Wevrick 2000). The ICR for this region, and where imprinting mutations occur, lies in a region at the 5' end of SNURF-SNRPN, a bicistronic transcript encoding a small protein (SNURF) and a subunit (SMN) of a ribonucleoprotein (Sutcliffe et al. 1994; Gray et al. 1999). Deletion of this ICR and associated CpG island causes loss of imprinting for the entire region, including SNURF-SNRPN, IPW (Wevrick et al. 1994), a collection of snoRNAs (Cavaille

et al. 2000; de los Santos et al. 2000), *NDN* (MacDonald and Wevrick 1997), *MAGEL2* (Lee et al. 2000), *MKRN3* (Jong et al. 1999), plus *Frat3* in mouse (Chai et al. 2001; Kobayashi et al. 2002) which are paternally expressed, and *UBE3A* (Rougeulle et al. 1997) and *ATP10C* (Meguro et al. 2001), which are maternally expressed (Figure 1-2).



Figure 1-2. Genomic organization of the human PWS/AS imprinted domain.

Chromosome 15 is shown at top, with the imprinted domain indicated by the red box. The 2.5 Mb domain is shown to scale in the center frame with transcripts indicated in blue. Note the large distance between the IC and the transposed imprinted genes. Genes are indicated at the bottom as a schematic. The IC is coincident with *SNRPN* exon 1.

The function of this ICR, more commonly referred to as the imprinting center (IC) at the PWS/AS locus, has been carefully dissected using a variety of patient studies, transgenic mice and other experiments. It was clear from UPD patients that PWS was due to a lack of paternally expressed genes instead of overexpression of maternally active genes from 15q11-13, since PWS resulting from maternal UPD had the same phenotype as paternal deletion (Glenn et al. 1996). A small number of cases are due to a heterogeneous class of imprinting mutations, either carrying microdeletions, which are often familial, and what are thought to be developmental or stochastic failures in the imprint process, which are sporadic (Nicholls et al. 1998). These microdeletions define an IC that lies at the 5' end of the SNRPN gene, where upstream exons not part of the protein coding potential of SNRPN have been detected (Figure 1-3) (Sutcliffe et al. 1994). These are referred to as the IC transcripts and they are important to the imprinting process as a splice mutation has been found in an AS patient (Dittrich et al. 1996). The microdeletion patients also delineate a bipartite functional structure for the PWS/AS IC, where a collection of AS patients define a smallest region of overlap (SRO), necessary for normal paternal to maternal allele identity switch, and a set of PWS patients define the PWS-SRO, necessary in the maternal to paternal switch (Figure 1-3) (Buiting et al. 1995; Saitoh et al. 1996). For example, a male inheriting a microdeletion of the PWS-SRO from his mother will not be able to reassign that allele to a paternal epigenotype for his progeny, who will have a 50% chance of inheriting this grandmaternal epigenotype from this male and develop PWS. A female on the other hand, inheriting the same microdeletion from her father will be able to reassign

the epigenotype to a maternal one. Male to male and female to female transmission does not require reassignment.



Figure 1-3. Upstream exons of SNURF-SNRPN and patient microdeletions.

At top is a schematic map (not to scale) of the upstream region of *SNURF-SNRPN*, with upstream exons. Nomenclature of these exons is provided below, with two previously published naming schemes. Bars below represent the extent of the microdeletions in patients that define the PWS (blue) and AS (red) SROs.

The region of conserved synteny in mouse chromosome 7C is likewise imprinted and behaves in much the same way. While useful in showing the conserved imprinted characteristics, the mouse models for the pathological phenotype were less clear. The PWS and AS models, achieved through maternal and paternal duplication of mouse 7C, did not recapitulate some aspects of the diseases (Cattanach et al. 1992; Cattanach et al. 1997). These were similar to a mouse strain carrying a fortuitous large deletion of the entire 7C imprinted domain (Gabriel et al. 1999). In retrospect, the difference in phenotype was not surprising, considering the number of genes involved and the potential that each may have subtle species-specific differences in function. However, the mouse system has been shown to be an excellent model for the study of imprinting in this region. Targeted deletions of part of Snrpn in a functional domain of the SmN protein showed that disruption of this gene had no effect on viability or imprinting, but a larger deletion involving the 5' end of Snrpn did have a phenotype (Yang et al. 1998). These mice shared the same failure to thrive phenotype as the maternal UPD mice (Cattanach et al. 1992). This phenotype is postulated to be causally related to the hypotonia seen in PWS neonates (Yang et al. 1998). More importantly, imprinting for the entire region was perturbed including dysregulation of imprinted genes more than a Megabase away.

How the PWS-SRO and the AS-SRO elements bring about the imprint switch for the locus was shown by recent work on the mechanisms by which these elements interact. While these two elements are 35kb apart endogenously, when brought together in close proximity, a transgenic construct is able to carry out all the steps of the imprinting process (Shemer et al. 2000). Using a series of transgenic mice with varying parts of the mouse

equivalent of the PWS-SRO and the human AS-SRO, it was found that one kilobase of the human AS-SRO and 200bp of the mouse *Snrpn* minimal promoter was sufficient for imprinting of a reporter gene. In addition to paternal specific expression, the transgene also showed differential DNA methylation and asynchronous replication, recapitulating these features of the endogenous locus. Further dissection of the step-wise function of this IC construct has shown that the mouse PWS-SRO contains elements necessary for *de novo* methylation of the maternal allele in imprint establishment and elements necessary for maintenance of the imprinted state in somatic tissues during development (Kantor et al. 2004a). In addition, there is an element that prevents methylation on the paternal allele. Therefore, the PWS-SRO is not simply a locus control region being controlled by the AS-SRO, but carries signals necessary and specific to imprinting. While the AS-SRO is upstream of the PWS-SRO, since the AS-SRO is not a simple matter that the AS-SRO imprints the PWS-SRO, since the AS-SRO is not able to imprint an unrelated β -globin locus element (Shemer et al. 2000).

The situation in the mouse is complicated by the observation that a small 0.9 kb deletion of the region homologous to the human PWS-SRO does not perturb imprinting (Bressler et al. 2001), whereas a larger 35 kb deletion does (Yang et al. 1998; Chamberlain and Brannan 2001), suggesting *Snrpn* exon1 and the associated CpG island are not important in imprinting in the mouse. The explanation was revealed when it was discovered that the mouse has a second redundant imprinting center capable of functioning in the absence of the first (Figure 1-4) (Kantor et al. 2004b). A 4.8 kb deletion that deletes the first IC and part of the second exhibits partial imprinting defects (Bressler et al. 2001).

How the remnants of the second IC are able to carry out partial imprinting is not well characterized.

•



Figure 1-4. Second functional IC in the mouse PWS/AS region.

Schematic of the two redundant ICs equivalent to the PWS-SRO in the mouse. IC1 is coincident with exon 1 of *Snurf/Snrpn*, while IC2 is in the first intron. Below are mouse deletions of the IC, and their effects on imprinting (+, imprinting retained, -, imprinting disrupted).



Figure 1-5. Interaction between PWS-SRO and AS-SRO. The PWS-SRO is believed to be a positive acting element on the imprinted domain, while the AS-SRO is believed to be a negative regulator of the PWS-SRO.

Further understanding of the cis-acting genetic epistasis between the two SRO elements was gained from studies using patient cells carrying microdeletions of the IC (Perk et al. 2002). It was found that the AS-SRO is associated with maternal allele-specific histone acetylation, H3 lysine 4 dimethylation (H3K4me2), and DNase I hypersensitivity, but surprisingly, not differential DNA methylation. In transgenic experiments however, differential DNA methylation are also observed (Kantor et al. 2004a). These epigenetic characteristics are not affected by deletion of the PWS-SRO in patients. This is in contrast to the reciprocal situation, whereby deletion of the AS-SRO causes biallelic hypomethylation, DNase I hypersensitivity and H3K4me2 modification at the PWS-SRO, features usually found only on the paternal allele, yet does not affect paternal allele-specific histone acetylation and asynchronous replication. From these observations, it is clear that the AS-SRO is necessary for the maternal methylation and certain other characteristics of the PWS-SRO, but does not depend on the PWS-SRO for its own epigenetic state. The PWS-SRO seems to be responsible for activation of paternal genes in cis, while the AS-SRO represses the PWS-SRO in *cis* on the maternal allele (Brannan and Bartolomei 1999) (Figure 1-5). This epistasis explains why deletion of both elements results only in PWS and not AS. When both are deleted, the paternal genes are not activated by the PWS-SRO on the paternal allele, while on the maternal allele, the AS-SRO is no longer required to repress the PWS-SRO.

As described, many genetic lesions lead to PWS and AS, but it is the epigenetic characteristics of this locus that are of interest here. The importance of epigenetic marking is demonstrated by a class of PWS and AS patients where no genetic lesions are found.

They are thought to carry epimutations, or errors in the epigenetic marking of alleles (Buiting et al. 2003). It is therefore critical to the understanding of PWS and AS imprinting to characterize the epigenetic regulation of the locus. In addition to DNA methylation, numerous lines of evidence indicate allele-specific differences in chromatin structure. Replication timing is often correlated to chromatin states in that euchromatin replicates earlier in S-phase than does heterochromatin (Goren and Cedar 2003). It has previously been observed that the two alleles of the PWS region replicate asynchronously (Knoll et al. 1994). Paternal-early and maternal-late replication was observed near the imprinted domain, while other patterns were observed more distally. DNase I hypersensitivity is commonly correlated with open chromatin (Weintraub and Groudine 1976). Two strong paternal allele-specific DNase I hypersensitivity sites flank exon 1 of SNRPN, while remaining resistant on the maternal allele (Schweizer et al. 1999). Interestingly, a less striking site of maternal allele-specific DNase I sensitivity was also observed coincident with the AS-SRO. Chromatin compaction as measured by density fractionation gave complementary results, where the paternal allele was measured to be less compact than the maternal allele (Watanabe et al. 2000). Larger scale organization has also been examined using two different fluorescence in situ hybridization (FISH) techniques, both indicating allele-specific patterns of matrix association (Greally et al. 1999; Kagotani et al. 2002).

The IC obviously plays an essential role at the PWS/AS imprinted domain. It is responsible for the correct expression of genes spread out over two Megabases, allelic identity and its maintenance throughout development. Certain details of how this genetic

element receives the initial epigenetic mark have been elucidated. How this translates into long range regulation of genes in *cis* however is less clear.

Long IC transcript model

The genomic organization of the imprinted region between the IC and the maternally expressed gene *UBE3A* includes more than the aforementioned paternally expressed *IPW* and snoRNA genes and the maternally expressed *ATP10C*. There are also other transcripts such as the *PAR* (for Prader-Willi/Angelman region) transcripts of unknown function (*PAR-1, PAR-2, PAR-4, PAR-5, PAR-7* (Sutcliffe et al. 1994), *PAR-SN* (Ning et al. 1996)), some of which have been shown to be paternally expressed, and only in certain tissues such as brain and skeletal muscle, in contrast to the ubiquitous expression of *SNURF/SNRPN*. In addition, there is also a paternally expressed transcript in antisense orientation to *UBE3A* (Rougeulle et al. 1998) that is also restricted in tissue-specific expression patterns and is under imprinted control of the IC (*UBE3A-AS*, (Chamberlain and Brannan 2001)).

Study of the IC has always been daunting because of its transcriptional complexity. The basic structure of the *SNURF/SNRPN* gene includes ten exons and is unusual in that it encodes a bicistronic transcript with two open reading frames, one for SMN (exons 4-10) and the SNRPN upstream reading frame protein (*SNURF*, exons 1-3) (Dittrich et al. 1996; Gray et al. 1999). However, operons are rare in mammalian genomes and the significance of this bicistronic gene to imprinting is unclear. While exon 1 is associated with a differentially methylated CpG island and is the most commonly transcribed 5' end to the *SNURF/SNRPN* gene, many 5' upstream exons exist and are thought to function in
imprinting (Dittrich et al. 1996; Farber et al. 1999). There are at least seven upstream exons in human, with several others that may be pseudo-exons (Figure 1-3). Most of these special U transcripts initiate at either u1B or u1A, exclude exon 1, splice directly to exon 2, and include variable numbers of the rest of the *SNRPN* exons (Farber et al. 1999). Most significantly, exon u5 is deleted in all AS patients with a microdeletion of the IC, suggesting this exon and perhaps upstream transcription plays a critical role in the paternal to maternal imprint switch. Also, whereas expression of *SNURF/SNRPN* is ubiquitous, transcripts containing these upstream exons are mainly in tissues such as brain, heart, testis and ovary (Dittrich et al. 1996). The significance of this will not become apparent until put into context of the rest of the transcription unit.

This transcription unit becomes more complex with addition of extra 3' exons into the picture. As with the 5' exons, each report provided evidence of more and more 3' exons. First were exons immediately downstream of *SNRPN* (10a, 11, 12) that connected to the *PAR* transcripts, previously thought to be individually expressed (Buiting et al. 1997), then to a larger set (13-20) that encompassed some of the snoRNAs (Wirth et al. 2001). It was also shown that some of these exons are only found in certain tissues that may not include *SNURF/SNRPN* exons (Buiting et al. 1997).

This transcriptional nightmare was brought into a single model with the realization that all of the paternally expressed transcripts between the IC upstream exons and *UBE3A-AS* represent alternative transcripts of a single transcription unit at least 460kb in size (Runte et al. 2001). This was first suggested by the fact that all the paternally expressed transcripts are transcribed from centromere to telomere whereas both maternal genes are transcribed in the opposite direction. Through the sequencing of multiple RT-PCR products and ESTs, a total of 128 new 3'exons were found distributed throughout the telomeric side of the human imprinted domain, linking pieces of previously known paternal transcripts. There was a high degree of alternative splicing for the products described, but many seemed to overlap the snoRNA genes, probably acting as host transcripts for these promoter-less intronic functional RNAs. The current model is that the paternal long transcript imparts paternal expression of genes in cis and in the same transcriptional direction, and downregulates the paternal allele of the maternally expressed gene UBE3A by an antisense mechanism since it is transcribed in the opposing direction (Figure 1-6). Another twist to the model was presented with evidence of the same long transcriptional unit in the mouse (Landers et al. 2004). In the mouse, it was observed that many of the alternative transcripts start with the upstream exons, some of which are 500kb upstream of Snrpn exon 1, making the transcription unit 1 Mb in length. Some of these transcripts also spliced in such a way as to exclude Snrpn. The most interesting lesson learned from the mouse is that these transcripts starting at the upstream exons and ending with Ube3a-as are tissue-specific, which provides an explanation as to how the imprinting of Ube3a is tissuespecific while Snurf/Snrpn is not. It is probable that this mechanism also operates in the human but this awaits experimental verification. How the maternal expression of ATP10C fits into this model is also not known, but it is possible further work will find exons extending telomeric to UBE3A.



Figure 1-6. Long paternal transcript through telomeric PWS/AS region.

The 148 identified exons part of the paternal transcript controlling imprinting of paternal and maternal genes telomeric to the IC. *HBII* genes are the human snoRNA genes. Reproduced from figure 5A, (Runte et al. 2001).

Transposed imprinted genes

While the long transcript model is a satisfying explanation for imprinting of the genes between the IC and ATP10C, and the tissue specificity of imprinting, it is not consistent with the imprinting of the paternally expressed genes NDN (MacDonald and Wevrick 1997), MAGEL2 (Lee et al. 2000), and MKRN3 (Jong et al. 1999) on the centromeric side of the IC (in human, telomeric side in mouse). The first inconsistency is the transcriptional direction of these genes, with MKRN3 transcribed from centromere to telomere and NDN and MAGEL2 transcribed in the opposite direction (Figure 1-2). In the mouse, an additional gene exists called Frat3, which is the product of a more recent transposition event (Chai et al. 2001; Kobayashi et al. 2002). All four genes seem to be processed forms of other genes, indicating they are likely products of past retrotransposition events. Therefore, they will be referred to as the transposed imprinted genes. All of these genes are associated with a differentially methylated region in their CpG islands, in contrast to the genes thought to be under control of the long antisense transcript. This fits well with the model that most maternally silenced genes are mediated by DNA methylation, and most paternally silenced genes are associated with an antisense mechanism (Reik and Walter 2001). A theory has been proposed to explain this difference that is based on the early demethylation events of the paternal genome in the zygote. Whereas the maternal genome demethylates passively, the paternal genome may have evolved other mechanisms such as antisense transcription to silence imprinted genes in spite of active demethylation (Mayer et al. 2000). Therefore, while the IC can silence the paternal alleles of the maternally expressed genes UBE3A and ATP10C via the long

antisense transcript, a separate mechanism may exist for regulation of the paternally expressed genes such as the transposed imprinted genes, which carry individual differentially methylated CpG islands. In light of the long transcript, it is clear how the action of the IC can span the 460kb from the IC to *UBE3A* in its influence. Outside of this model, there is no proven way for the IC to influence the transposed imprinted genes at a distance. Characterization of the mechanisms of imprinting for these genes will be critical in the understanding of the function of the IC. This will be explored in detail in the following chapters. To understand the regulation of the transposed imprinted genes, it will be useful to study one as a model for the others in the cluster. Chapter 3 will focus on the immediate regulation of the promoter of *NDN*. Chapter 4 builds on the observations at the promoter by examining the chromatin context of the region containing *NDN*. Chapter 5 investigates the mechanism of the entire transposed imprinted domain and will reveal a possible model to reconcile the epigenetic regulation of imprinting and tissue-specific expression.

Epigenetics and Chromatin Control

The differential allelic regulation of imprinted genes is transmitted between generations and cell divisions without changes in DNA sequence. Memory of the parental origin and allelic identity is instead recorded in an epigenetic manner. DNA methylation, histone modifications and their interaction with each other are all involved in the orchestra of events that leads to correct imprinted gene regulation.

DNA methylation

Without a doubt, DNA methylation is the best studied epigenetic mark. It is truly an archetype of an epigenetic mark in that it sits atop DNA to carry extra information and it has a clear mechanism of heritability through its maintenance during DNA replication. It was not until the late 1970's when this minor base variant was associated with gene activity (Razin and Riggs 1980). Since then, its importance has been demonstrated in multiple systems such as tissue-specific regulation, differentiation, cancer, X-inactivation, and genomic imprinting (Paulsen and Ferguson-Smith 2001). Methylation of CpG dinucleotides of mammalian genomes is accomplished by DNA methyltransferases (DNMTs) that act on hemi-methylated DNA for maintenance (DNMT1) and Dnmts that act on unmethylated DNA to generate *de novo* methylation patterns (DNMT3A/B) (Bestor 2000). Mutations in *DNMT3B* are found in ICF syndrome, which shows immunodeficiency and centromeric instability on certain chromosomes (OMIM: 242860, (Hansen et al. 1999; Xu et al. 1999)). Targeted deletions of the *Dnmts* are lethal, with *Dnmt1*-nulls being the most severe, and affecting imprinted genes and X-inactivation (Li et al. 1992; Li et al.

1993; Beard et al. 1995; Okano et al. 1999). Over-expression of *Dnmt1* causes hypermethylation and loss of imprinting (Biniszkiewicz et al. 2002). Other DNMTs such as DNMT2 and DNMT3L exist, though they are without *in vitro* methylating activities, and may participate in pathways related to their true methyltransferase relatives (Bestor 2000).

DNA methylation is thought to modify protein-DNA interactions to bring about its biological effects (Razin and Riggs 1980). Many transcription factors are not able to bind if their target DNA is methylated (Tate and Bird 1993). The DNA binding factor CTCF has received much attention of late and has been shown to be sensitive to DNA methylation (Bell and Felsenfeld 2000; Hark et al. 2000). This will be discussed in a following section (page 54). DNA methylation does not always preclude binding of factors, and in fact, there are protein domains with specific affinity for methylated DNA (Hendrich and Tweedie 2003). There are five proteins with a methyl CpG binding domain (MBD); MeCP2, MBD1, MBD2, MBD3 and MBD4. Most have been shown to have a repressive activity, while MBD4 seems to function in a repair pathway that deals with the mutagenic cost of having methylated cytosine residues in the genome (Millar et al. 2002). MBD2 may also have direct DNA demethylase activity but this has been controversial (Bhattacharya et al. 1999; Ng et al. 1999). These proteins bring about transcriptional repression through recruitment of other proteins such as chromatin remodeling enzymes, histone deacetylases (HDACs) and histone methyltransferases (HMTs). MBD2 and MBD3 for example, are associated with the MeCP1 and NuRD repressive complexes respectively (Ng et al. 1999; Zhang et al. 1999). MeCP2 in particular has been associated with HDAC activities (Nan et al. 1998) as well as HMT activities (Fuks et al. 2003), and even associates with DNMT1 (Kimura and

Shiota 2003). Beyond that, much attention has been paid to *MECP2*, because it is a causative gene in Rett Syndrome (Bienvenu et al. 2000). Although widely expressed and thought to have a general role in gene repression, phenotypic defects are mainly neurological, perhaps due to sensitivity of the brain to loss of MeCP2 and perturbation in a subset of genes it regulates (Chen et al. 2001; Guy et al. 2001; Tudor et al. 2002). Most importantly to our discussion, there are isolated examples of loss of imprinting (LOI) in Rett Syndrome (Horike et al. 2005; Makedonski et al. 2005), although there are no obvious global defects in imprinting (Balmer et al. 2002).

Through the action of the maintenance methylase on newly synthesized DNA, methylation patterns can be transmitted through cell divisions. But in terms of imprinting, epigenetic signals must be transmitted through generations to record the parental gender of origin of an allele. This presents a challenge for DNA methylation as there is a global demethylation event early in zygotic development (Jaenisch 1997; Morgan et al. 2005). This challenge is especially daunting for the paternal genome since it is actively demethylated in the zygote after fertilization and before the first round of DNA replication. Fortunately, DMRs associated with ICs are able to retain their methylation state through an unknown mechanism and represent *bona fide* imprints from the last generation. However, many DMRs not part of ICs lose their methylation and must be re-established after the demethylation event (Hanel and Wevrick 2001). To combat the active demethylation of the paternal genome by the oocyte in the zygote, an additional mechanism may have evolved (Mayer et al. 2000). It has been suggested that the preponderance of antisense transcripts seen for imprinted genes that act to suppress the paternal allele is to replace the repressive

activity of DNA methylation, and is therefore another weapon in the sexual conflict theory of imprinting.

Although it is widely held that DNA methylation is important to mammalian development (Eden and Cedar 1994; Meehan 2003), it is not without debate (Walsh and Bestor 1999). The vast majority of methylated DNA in the mammalian genome is found at transposable repetitive elements (Yoder et al. 1997). It has been proposed that this is the true function of DNA methylation, as host defense against transposition and that its use in imprinting and X-inactivation may be secondary and may not be important for development otherwise (Bestor 2000). This is still controversial and the fact that transpositions are still controlled during the stages of demethylation argues against this (Bird 1997). Others have suggested DNA methylation is necessary to control background transcriptional noise (Bird 1997) or to modify the regulatory effects of transposed elements indirectly (Martienssen 1998). Whatever the evolutionary function of DNA methylation, it clearly has a major role in imprinted gene regulation.

Histone modifications

Nucleosomes are the basic monomeric unit of chromatin. With an amazing capacity to compact two meters of DNA into a eukaryotic nucleus, it was easy to assume the nucleosome's function was one of simple compression. Of course it is now known that histones plays an important role in gene regulation (Strahl and Allis 2000; Felsenfeld and Groudine 2003). The basic structure of the nucleosome consists of two each of the four core histone subunits H4, H3, H2A and H2B, with approximately 146 bp of DNA coiled around a basic groove in the complex. As a polymer, nucleosomes are central to higher

order chromatin organization (Figure 1-7), especially in the 11 nm and 30 nm configurations where the unstructured histone tail domains are exposed and accessible to modification.



Figure 1-7. Levels of chromatin compaction.

Sequential compaction of DNA allows compression as well as regulation at different levels. Reproduced from Figure 1A of (Felsenfeld and Groudine 2003).

Despite its status as darling of the epigenetics field in recent years, histone modification as it pertains to gene regulation is not a new idea. Histone acetylation was associated with gene activation and transcription almost forty years ago (Pogo et al. 1966), but of course without the detailed *in vivo* characterization that was made possible by chromatin immunoprecipitation (ChIP) (Kuo and Allis 1999). Perhaps more crucially, antibody reagents capable of recognizing specifically modified histones in ChIP were developed (Hebbes et al. 1988). The ChIP assay represented for the first time a method by which protein-DNA interactions in vivo can be studied for any endogenous locus in a sequence-specific manner. Since then, the field advanced as fast as new antibody specificities could be produced and characterized. A new nomenclature evolved to deal with this new language (Turner 2005). For simplicity, the rest of this text will use this nomenclature convention for discussion involving histone modifications (Table 1-1). With each new report detailing association between a chromatin-templated biological process and a specific histone modification, it became clear this represented a novel level of information in gene regulation (Figure 1-8, Table 1-2). The histone code hypothesis was proposed, which suggests that combinations of multiple covalent modifications on histones specify unique biological function (Strahl and Allis 2000; Grant 2001; Iizuka and Smith 2003; Peterson and Laniel 2004; Dion et al. 2005).

Modifying group	Amino acid(s) modified	Level of modification	Abbreviation for modification	Examples of modified residues
Acetyl-	Lysine	mono-	ac	H3K9ac
Methyl-	Arginine Arginine	mono- di-, symmetrical	mel me2s	H3R17me1 H3R2me2s
	Arginine	di-, asymmetrical	me2a	H3R17me2a
	Lysine	mono-	me1	H3K4me1
	Lysine	di-	me2	H3K4me2
	Lysine	tri-	me3	H3K4me3
Phosphoryl-	Serine or threonine	mono-	ph	H3S10ph
Ubiquityl-	Lysine	mono-	ubl	H2BK123ub1
SUMOyl-	Lysine	mono-	รม	H4K5su
ADP ribosyl-	Glutamate	mono-	arl	H2BE2ar1
	Glutamate	poly-	arn	H2BE2arn

Table 1-1. Brno nomenclature for histone modifications.

Abbreviation scheme for histone modifications, with subunit first, then residue, and modification. Named after Brno, Czech Republic, where the nomenclature was proposed. Reproduced from Table 1 of (Turner 2005).



Figure 1-8. Covalent modifications of canonical histone subunit tails. All four histone tails can carry covalent modifications. Ac = acetylation, P = phosphorylation, Me = methylation, Ub = ubiquitination. Reproduced from Figure 1 of (Peterson and Laniel 2004).

Modification	Histone	Site	Enzyme	Possible function
Acetylation	H2A	K4 (S. cerevisiao)	Esal	Transcriptional activation
		K5 (mammals)	Tip60	Transcriptional activation
			p300/CBP	Transcriptional activation
		K7 (S. cerevisize)	Hatt	?
			Esal	Transcriptional activation
	H2B	К5	ATF2	Transcriptional activation
		K11 (S. cerevisiae)	Gen5	Transcriptional activation
		K12 (mammals)	p300/CBP	Transcriptional activation
			ATF2	Transcriptional activation
		K16 (S. cerevisiae)	Gcn5	Transcriptional activation
			Esa1	• • • • • • • • • • • • • • • • • • • •
		K15 (mammals)	p300/CBP	
			ATE2	Transcriptional activation
		K20	p300	Transcriptional activation
	U 2	KA	Eeo1	Transcriptional activation
			Loui Lino?	2
		KO	2	f History deposition
		N9	f CT	
			6675	Transcriptional activation
			SHC-1	Transcriptional activation
		K14	Gcn5, PCAF	Transcriptional activation
			Esal, Tip60	Transcriptional activation
				DNA repair
			SRC-1	Transcriptional activation
			Elp3	Transcription elongation
			Hpa2	?
			hTFIIIC90	RNA polymerase III transcription
			TAF1	RNA polymerase II transcription
			Sas2	Euchromatin?
			Sas3	Transcriptional activation/elongation?
			p300	Transcriptional activation
		K18	Gcn5 (SAGA/STAGA complex)	Transcriptional activation
				DNA repair
			p300, CBP	DNA replication
				Transcriptional activation
	e de la composición d	K23	Gcn5 (SAGA/STAGA complex)	Transcriptional activation
			Sas3	DNA repair
		an an tao ang sa s	p300, CBP	Transcriptional activation/elongation?
				Transcriptional activation
		K27	Gcn5	Transcriptional activation
	на	К5	Hari	Histope deposition
			Feat. Tip60	Transcriptional activation
				DNA repair
			ATE2	Transcriptional activation
			цтэ)	a
			0300	Transcriptional activation
		K8	Gen5 PCAF	Transcriptional activation
	• • • • • • • • • • • •		Sent Tinff)	Transcriptional activation
	ling of the		Louis libos	
			ATE0	Transportional active for
			A1F2	
			Elp3	Iranscription elongation
para di Ang	en por l'Alia		p300	Transcriptional activation
	ante stato di	K12	Hatt	Histone deposition
				Telomeric silencing
			Esa1, Tip60	Transcriptional activation
Aller and Aller		an an the storage fitting		DNA repair

Table 1-2. Summary of modifications.

Table of modifications, their modifiers and functions. Reproduced from (Peterson and Laniel 2004).

Modification	Histone	Site	Enzyme	Possible Function
		K12	Hpa2	?
		K16	Gcn5	Transcriptional activation
			MOF (D. melanogaster)	Transcriptional activation
				Transcriptional activation
			Esa1 (veast), Tip60 (mammals)	DNA repair
			ATF2	Transcriptional activation
			Sas2	Euchromatin
Methylation	H3	KA	Set1 (vnast)	Permissive exchromatin (di-Me)
			Set9 (vertebrates)	Active euchromatin (tri-Me)
				Transcriptional elongation/memory (tri-Me)
				Transcriptional activation
			MIL Tor	Transcriptional activation
			Achi (D. malanagastad	Transcriptional activation
		Vo	Asiri (D, molanogasion)	Transcriptional activation
		K9	SDASAIT CR4	DNA methodoline (million)
				DNA methylauon (tn-me)
			623	I ranscriptional repression
			SETDRI	Transmitting
			Dim 6 Kamtarita	DNA methylation (tri \$5.)
			Dim-5, Krypionite	DNA metrylation (un-wie)
		D / 7	Asn1 (D, melanogaster)	Transcriptional activation
		H1/	CARMI	I ranscriptional activation
		K27	E202	I ranscriptional silencing
				X inactivation (tn-Me)
		K36	Set2	Transcriptional elongation
		· · · · · · · · · · · · · · · · · · ·		Transcriptional repression?
		K79	Dot1p	Euchromatin
				Transcriptional elongation / memory
	H4	R3	PRMTI	Transcriptional activation
		K20	PR-Set7	Transcriptional silencing (mono-Me)
			Suv4-20h	Heterochromatin (tri-Me)
			Ash1 (D. melanogaster)	Transcriptional activation
		K59	?	Transcriptional silencing?
Phosphorylation	H2A	S1	• ?	Mitosis
			7	Chromatin assembly?
			MSK1	Transcriptional repression
		T119	NHK-1	Mitosis
		S129 (S. cerevisiae)	Mec1	DNA repair
		S139 (mammalian H2AX)	ATR, ATM, DNA-PK	DNA repair
	H2B	S14 (vertebrates)	Mst1	Apoptosis
	and the set	S33 (D. melanogaster)	TAF1	Transcriptional activation
	H3	73	a , 1997 - 199	Mitosis
		\$10	Aurora-B kinase	Mitosis, meiosis
			MSK1.MSK2	Immediate-early activation
			Snf1	Transcriptional activation
		T11 (mammals)	Dik/ZIP	Mitosis
		S28 (mammals)	Aurora-B kinase?	Mitosis
Media de la composición de la composicinde la composición de la composición de la composición de la co		- To functional of	MSK1. MSK2	Immediate-early activation
	На	St	?	Mitosis
Hispitetos	H2A	K110 (mammale)	HR64 B7	Spormatogenetic
opiquiquation	1000	Kt 00 (momental)	LIDEA D2	Algoria Maloria
	п <i>с</i> р	Ki 20 (mainmais)	Dedf	
and the day	and a second	NI 23 (3. Cerevisiae)	006m	ranscriptional activation
				Euchromatin
	H3	?	?	Spermatogenesis
Sumoviation	HA	7	Libes	Transcriptional repression

Table 1-2, continued. Summary of modifications.

Table of modifications, their modifiers and functions. Reproduced from (Peterson and Laniel 2004).

Histones are also one of very few candidates for carrying epigenetic information. It is clear how DNA methylation is transmitted past DNA replication, and there are models to suggest how histone modifications can be transmitted. Most importantly, it has been shown that nucleosomes are retained on daughter strands past the replication fork (Figure 1-9 A) (Bonne-Andrea et al. 1990), and it is thought that parental histones are distributed equally to newly synthesized DNA with the addition of new nucleosomes (Krude 1999; Lucchini et al. 2001). This dilution of parental histories and their covalent modifications is strikingly analogous to the hemi-methylated state of newly synthesized DNA. Spread and propagation of the modification state of parental nucleosomes to new nucleosomes can then accomplish maintenance of these states through cell divisions. This is still controversial however as some sites of modification thought to be important in gene regulation span only several histones, raising the doubt that faithful distribution to daughter strands is possible (Henikoff et al. 2004). A model by which histone halves are split off to daughter strands, while attractive in terms of epigenetic transmission, has very little experimental support (Figure 1-9 B). These states are obviously propagated though, and in some cases without DNA methylation, so it is likely a matter of when, instead of if, these mechanisms will be found. In fact, an alternate model of epigenetic inheritance involving variant histones will be discussed later (Figure 1-10).



Figure 1-9. Models of histone conservation during replication.

(A) Nondispersive model where old nucleosomes (dark brown) are segregated randomly to leading and lagging strands past the replication fork (green pentagon) with new nucleosomes (light brown). (B) Semiconservative model where new subunits are incorporated with old subunits. In this model, it is one pair of H3-H4 and H2A-H2B that is conserved with new subunits.

As with DNA methylation, there are proteins able to read, write and erase the information on histories. But unlike the biochemical simplicity of DNA methylation, histone modification is highly variable and requires families of enzymes, and variants of domains to carry out downstream biological directives. Many of these enzymes have previously been found to be transcriptional co-activators or co-repressors (Grant 2001). Histone acetylation has been most well characterized (Grunstein 1997). It is associated with gene activity as well as potentiation of transcription (Hebbes et al. 1988). Families of histone acetyltransferases (HATs) and histone deacetylases (HDACs) catalyze this modification on lysine residues on all four subunits of the nucleosome. Many HATs and HDACs have been characterized, some with preferences for certain residues. Two types of HATs exist: nuclear type-A, involved in gene regulation, and cytoplasmic type-B, involved in pre-acetylation of newly synthesized histones for assembly. GCN5 is one of the first HATs characterized in yeast and it is required for activation of many genes (Kuo et al. 1996). Other HATs such as CBP and p300 have been shown to be involved in gene activation in response to cellular signals (Chakravarti et al. 1996). They also do not act alone, but participate in large complexes that are able to target their activities. Some are also able to acetylate other proteins, often contributing to gene activation. Similarly, HDACs are part of large repressive complexes and are recruited to genes through complex interactions (Grunstein 1997). While the acetylation of histones can bring about changes in chromatin structure by neutralizing the positive charge that facilitates interaction with DNA, acetylation can also act through signal transduction by creating binding sites for the bromodomain (de la Cruz et al. 2005; Dion et al. 2005). Analogous to the function of the

MBD in the recognition of DNA methylation, bromodomains bind acetylated histones and are found in HATs, chromatin remodeling enzymes and HMTs, indicating their importance in chromatin and gene regulation. The observation that HATs contain bromodomains may be a significant indicator of how these modifications can be amplified or propagated to nearby residues, adjacent histone subunit tails, as well as neighboring nucleosomes.

In contrast to the harmonious picture of how histone acetylation functions in transcriptional regulation, histone phosphorylation seems to be involved in many different pathways. Phosphorylation was first linked to mitosis (Hsu et al. 2000; Nowak and Corces 2004), but has also been associated with transcriptional control (Peterson and Laniel 2004). Again, this may be through changes in charge of the nucleosomal particle or binding of specific factors, or most intriguingly, through interactions with other modifications, which will be discussed later. Recently, H2BS14ph has been shown to be involved in apoptosis by a specific kinase Mst1 (Ahn et al. 2005). Phosphorylation of histone variant H2A.X during damage has been shown to have an important role in DNA double strand break repair by INO80 complex recruitment (Morrison et al. 2004). Covalent addition of ubiquitin and SUMO groups has also been observed on histones and is linked to various chromatin functions. Unlike the poly-ubiquitination that is associated with protein turnover, histones are often mono-ubiquitinated, which may be related to mitosis, meiosis (Robzyk et al. 2000), transcription (Davie and Murphy 1994), and most importantly, to spermatogenesis (Jason et al. 2002). Sumoylation has been discovered recently and is associated with repression (Shiio and Eisenman 2003). ADP-ribosylation of histones is not well understood, but there is evidence this modification of the linker histone H1 may be

involved in memory formation (Cohen-Armon et al. 2004). While these processes are fascinating in their own right, this discussion will be restricted to modifications most relevant to transcriptional regulation.

Histone methylation has taken the limelight in recent years. While most histone acetylation states are associated with transcriptional activation, histone methylation can serve many roles, including activation and repression of gene activity (Peterson and Laniel 2004). Histone tails contain arginine (R) and lysine (K) residues that can be methylated. Furthermore, R and K residues can accommodate up to two and three methyl groups respectively, to elaborate the signal. In addition, dimethylation of R can be in two steric variants (Bannister et al. 2002). It is also believed that histone methylation has a longer half-life than acetylation or phosphorylation, making it a better candidate for long term memory of gene activity (Bannister et al. 2002). The first example was the characterization of CARM1 that is able to methylate H3 R residues and participates in the activation of genes under control of nuclear hormone receptors (Chen et al. 1999). It was soon found that H3K4me was a well conserved modification that was involved in gene activity (Strahl et al. 1999). Different levels of methylation at this lysine residue also correlated with different levels of activity; in this case, H3K4me2 and H3K4me3 with euchromatin and transcriptionally active genes, respectively (Santos-Rosa et al. 2002). H3K79me has been associated with telomeric silencing in yeast (Lacoste et al. 2002) and gene activation in yeast and mammals (Ng et al. 2003). In contrast, methylation of H3K9 is involved in heterochromatin assembly (Nakayama et al. 2001). This modification can be made by SUV39H1, the human homolog of a Drosophila gene long known to be involved in gene

regulation and position effects (Rea et al. 2000). SUV39H1 and other histone HMTs that act on lysines contain an evolutionarily conserved SET (Su(var), E(z), trithorax) domain, found in many proteins involved in transcriptional regulation (Xiao et al. 2003). While SUV39H1 is involved in the H3K9me of heterochromatin, another HMT, G9a is involved in the H3K9 and K27 methylation at euchromatic sites, necessary in transcriptional repression of developmental genes essential in embryogenesis (Tachibana et al. 2002; Roopra et al. 2004). Interestingly, G9a is essential in imprinting of the PWS/AS IC (Xin et al. 2003). Similar to bromodomain bearing proteins that carry out downstream effects of histone acetylation, specific proteins recognize histone methylation. H3K4me, a mark of activity, is able to disrupt binding of the NuRD repressor complex (Nishioka et al. 2002; Zegerman et al. 2002). H3K9me3, a mark of silencing, can be bound by the chromodomain of HP1, a major facultative heterochromatic protein (Bannister et al. 2001; Lachner et al. 2001). H3K27me however, is bound by polycomb group proteins (PcG) in regulation of developmental gene clusters (Kirmizis et al. 2004). H3K9me and H3K27me represent extremely similar epitopes, as evidenced by some antibodies that cannot distinguish the two, but they are uniquely identified by HP1 and PcG, illustrating the biological specificity of these marks (Fischle et al. 2003c). Subtle adjustment of H3K9 and K27 methylation levels are associated with facultative and constitutive heterochromatin with H3K9me3 and H3K27me1 at pericentric heterochromatin and H3K9me1/2 and H3K27me1/2/3 at silent euchromatin. With the number of residues and variation of methylation of each residue, a complex code can be elucidated from just histone methylation (Craig 2005). Recent advances have also shown that the histone methylation mark can truly be regulated by

erasure instead of the histone replacement during replication (Shi et al. 2004). Much like HDACs, LSD1 was identified as a co-repressor that specifically demethylates the H3K4 residue. It remains to be seen whether other histone methylation marks such as H3K9 and H3K27 can also be demethylated by a similar enzyme, but discovery of LSD1 will facilitate this search. It is also important to remember that while many modifying enzymes have been termed histone-specific, their substrates may not be so restricted, and HATs, HDACs, and HMTs may modify other proteins, often also involved in gene regulation (Chen et al. 1999; Robzyk et al. 2000; Girdwood et al. 2003).

Another way to reverse the effects of histone methylation or any other modification is the replacement of that histone, either during DNA replication or by replication independent means (Ahmad and Henikoff 2002). While canonical histone H3 subunits are only synthesized and incorporated during replication, variants can be added at other times, and by special complexes. H3.3 is a special H3 replacement variant that differs by only four amino acids and is incorporated into heavily transcribed euchromatin (Tagami et al. 2004). Even variants are modified as canonical histones are. H3.3 is often enriched in active modifications (McKittrick et al. 2004). It has been proposed that epigenetic inheritance of chromatin states can be achieved through the use of H3.3 (Henikoff et al. 2004). An actively transcribed region would be enriched in H3.3 which, after dilution to daughter strands during replication would direct the transcription of the locus in the daughter cell, thereby enriching the region again for H3.3 (Figure 1-10). In this scheme, it is transcription that takes on the role of maintenance, versus amplification of histone modifications having this role in the previous model of transmission of histone states. Note

that of course, these are not mutually exclusive models. It is also suggested that this is how the activation states of certain regulatory elements are transmitted through intergenic transcription (Rank et al. 2002). It should be noted here, that this model is distinct from antisense regulation which inactivates genes by transcription in *cis*.



Figure 1-10. Mechanism of transmission of H3.3 states through replication.

Transcription causes local replacement with variant H3.3. Replication would lead to a dilution of H3.3, but still leads to transcription of the locus, causing more replacement with H3.3, completing the cycle.

Adding yet another level of complexity is the central tenet of the histone code hypothesis: that different modifications act in sequence or in combinations to bring about unique biological properties (Strahl and Allis 2000). The most obvious example is the physical occupancy of a residue by a modification, thereby preventing further reactions. For example, the methylation of H3K9 would preclude acetylation at the same site. More commonly, examples have shown interactions between different residues (Figure 1-11) (Fischle et al. 2003b). H3S10ph has been shown to be a prerequisite for and to promote H3K14ac (Cheung et al. 2000; Lo et al. 2000). H3S10ph however, can be inhibited by methylation of the adjacent residue to H3K9me (Rea et al. 2000). To bring these interactions full circle, H3K9me depends on H3K14 deacetylation (Nakayama et al. 2001). This interactive and synergistic cycle may specify an ON and OFF state (Figure 1-12) (Berger 2001); other such "binary switches" have been characterized (Fischle et al. 2003a). Interactions can also be extremely specific: for example, H3K4me by Set9 inhibits Suv39h1 but not G9a mediated H3K9me (Nishioka et al. 2002). Modification on different tails can also affect each other. The best example is the unidirectional requirement of H2BK123ub for H3K4me, where a mutation of H2B K123R abolishes H3K4me but H3 K4R does not affect H2BK123ub levels (Sun and Allis 2002). One question comes to mind in consideration of these complex networks. Why must there be so many marks seemingly contributing to the same outcome? Perhaps slight adjustment of the code allows finer tuning than the simplicity of DNA methylation allows. The fluidity of histone modifications and the number of modifying enzymes in many pathways may also represent a point of integration of numerous cellular signals in gene regulation.

H3	(M)K4	(M)K9	Ac K9	PS10	AcK14	(M)K14 (M)R17 AcK1	B Ac K23	(M)R26	(M)K27	PS28	(M)K36 (M)K79
(M)K4		۵.	۲		0			Ø 1				
(M)K9	•		•	C			•	٢				
Ac K9	0	۵.										
PS10		٢	8		0							
AcK14/18/	23	¢										-
H4	PS1	(M)R3	Ac K5	Ac K8	Ac K12	Ac K16 (M)K20					
(M)R3				0								
Ac K5		0										
AcK8		©										
Ac K12		•										· · · · · · ·
Ac K16		•					•					Permissive
(M)K20						•	1 . 				6	Repressive

Figure 1-11. Examples of interactions between modifications.

Examples of permissive and inhibitory interactions between modifications on the histone tail on further modification of the same tail. Reproduced from Figure 1 of (Fischle et al. 2003b)



Figure 1-12. Binary switch theory.

Example of a binary switch of gene transcription, where one set of modifications synergistically has the opposite action of different set of modifications. Certain individual modifications may also have repressive effects on others.

Epigenetic interaction

An epigenetic hierarchy has often been sought, most importantly, between DNA methylation and histone modifications. Which is upstream in the pathway of gene regulation? As previously described, methylated DNA can recruit histone deacetylase activities through MBD containing proteins and complexes involved in repression (Bird and Wolffe 1999). Some proteins have also been found to contain a SET domain and a putative MBD, raising the possibility that DNA methylation can also direct histone methylation (Figure 1-13) (Zhang and Reinberg 2001). With the well established role of DNA methylation in gene regulation and its clear mode of inheritance, it was easy to speculate that it was higher up in the epigenetic hierarchy. Contrary to this, it was shown that in Neurospora, DNA methylation requires H3K9me. It was discovered that a mutation that causes decrease in DNA methylation (dim-5) was a HMT (Tamaru and Selker 2001). A similar situation was found in plants which carry CpNpG methylation as well as CpG methylation. Mutations in a plant HMT kyp abolished CpNpG methylation and H3K9me, but mutation of the CpNpG methylase *cmt3* only affected DNA methylation and not H3K9me (Johnson et al. 2002). It was later found that CMT3 bound, and required both H3K9me and H3K27me together to direct CpNpG methylation, showing the mechanism by which histone modifications can direct DNA methylation and adding to the histone code for silencing (Lindroth et al. 2004). Interestingly, histone modifications are more conserved across eukaryotes than DNA methylation, which is absent or negligible in organisms such as S. pombe and Drosophila (Nakayama et al. 2001). It has been proposed that DNA methylation is a more recent addition to the epigenetic schema to allow elaboration and

perhaps more stable marking of expression (Rice and Allis 2001), or as mentioned earlier, as a protective agent against genomic parasites (Bestor 2000). This is supported by the observation that the *cmt3* and *kyp* mutants show reactivation of certain retrotransposons in *Arabidopsis* (Bartee et al. 2001).



Figure 1-13. Interaction between DNA methylation and histone modification.

Examples of how DNA methylation and histone modifications can interact with each other. Repressive MBD proteins can bind methylated DNA (lollipop) and recruit HDAC activities to deacetylate histones. More directly, a protein can have both a MBD, as well as a histone modification domain such as a SET domain to methylate histones. Alternatively, a protein can bind methylated histones through a chromodomain and methylate DNA.

There are also important connections between gene regulation, histone modifications, DNA methylation and RNAi (RNA interference) (Matzke and Birchler 2005). RNAi is involved in aspects of transcriptional control at the RNA and chromatin level through the use of short homologous RNA molecules. Based on studies in plants, it has been found to be able to destroy specific mRNA and cause epigenetic alterations of DNA targets, but the existence of this pathway in mammals is still in question. As my work does not deal with RNAi and the links between its mechanisms and those proposed for antisense regulation of mammalian imprinted genes and X-inactivation are tenuous, this is outside the scope of this introduction.

Epigenetic mechanisms in genomic imprinting

The epigenetic hierarchy is especially pertinent to imprinting and X-inactivation. What comes first and what gets transmitted is central to the question of how two alleles in the same cell can maintain differential expression and epigenetic states independently of each other. With the mechanisms of transmission of histone modifications uncharacterized, DNA methylation is still the prime candidate for the imprinted mark, especially between generations, with the observation that ICs can carry differential methylation from the gametes (Soejima and Wagstaff 2005). However, the histone variant CENP-A is not replaced by protamines during spermatogenesis and is thought to be the mechanism by which centromeres are epigenetically transmitted (Henikoff et al. 2004), opening the possibility that a small subset of histones or their variants can also be retained. Histone variants are also correlated with the active and inactive X's (macroH2A and H2A-Bbd) (Chadwick and Willard 2001; Chadwick and Willard 2002). The relationship so far

between DNA methylation, histone modification and imprinting has varied by report, and no unifying theory exists. It is known that in *Dnmt1* null embryos, imprinting is perturbed for some but not other genes (Caspary et al. 1998; Xin et al. 2003). Acetylation was affected for H4 but not H3 in treatments with a DNA methylation inhibitor at SNRPN in human cell culture and resulted in reactivation of the silent maternal allele (Saitoh and Wada 2000). In mouse cell culture however, neither treatment with a HDAC inhibitor nor with the DNA methylation inhibitor was able to reactivate the silent allele of *Snrpn*, although some other imprinted genes were reactivated, and in some cases, in a heritable fashion after withdrawal of inhibitors (El Kharroubi et al. 2001). In G9a-null embryos, there is a loss of imprinting for Snrpn (Xin et al. 2003). This is in contrast to the Dnmt1 null that exhibits loss of differential DNA methylation at the IC, but retains differential H3K9me and imprinting. H3K9me is also one of the earliest events in X-inactivation after Xist coating (Heard et al. 2001; Mermoud et al. 2002). In a report studying which of DNA methylation, histone modifications, or antisense transcription was more correlated with tissue specific imprinting of IGF2R/Igf2R in human and mouse, it was found that H3K4me and H3K9me were most consistent with imprinted expression (Vu et al. 2004). Imprinting of the Kcnql imprinted domain in the placenta, which does not involve differential DNA methylation, is instead associated with H3K27me3, H3K9me2, and the PcG complex Eed-Ezh2 (Umlauf et al. 2004). The same PcG complex and H3K27me is also involved in Xinactivation (Plath et al. 2003), further implicating the evolutionary commonality between imprinting and X-inactivation (Lee 2003). Therefore, it is obvious that genomic imprinting and X-inactivation employ multiple epigenetic mechanisms in mono-allelic gene

regulation, but the players at various loci, tissues and species differ. Further examination of mechanisms involved will be instrumental in deciphering the language necessary for imprinting.

CTCF Biology

CTCF, the mammalian insulator protein

CTCF was first named for its ability to bind a CCCTC DNA sequence (Lobanenkov et al. 1990). But in fact, it is able to bind an array of different sequences in many species (Figure 1-14) (Ohlsson et al. 2001). It is highly conserved across vertebrates, and contains 11 zinc finger (ZF) domains, which gives it the ability to bind divergent sequences through combinatorial use of these domains (Filippova et al. 1996). Unlike other zinc finger proteins, the ZFs in CTCF are able to interact with proteins as well as DNA (Ohlsson et al. 2001). CTCF is able to activate or inhibit transcription in different situations depending on interactions with other proteins (Baniahmad et al. 1990; Awad et al. 1999). It is also able to recruit HDACs, linking it to histone modification (Lutz et al. 2000). Perhaps its most important function is at mammalian insulators. While the mechanism of insulator function is unknown (Bulger and Groudine 1999), it is thought that most if not all mammalian insulators bind CTCF (Bell et al. 1999). As mentioned earlier, CTCF function at an allelic insulator has a pivotal role in *Igf2/H19* imprinting (Bell and Felsenfeld 2000; Hark et al. 2000).



Figure 1-14. Variation in ZF domain usage in DNA recognition.

Chart showing guanine residues important in a subset of CTCF binding sites on the left. On the right, are the corresponding ZF domains used to contact those guanines, with white being most important and red dispensable in deleted CTCF constructs (pink being incomplete loss of binding when deleted). DMD4 and DMD7 are from the *Igf2/H19* ICR. Reproduced from Figure 2 of (Ohlsson et al. 2001).

One of the most well characterized CTCF binding insulators is the locus control region (LCR) of the β -globin genes (Bell et al. 1999). The β -globin LCR in many ways was at the forefront of the field of epigenetics. It is able to confer tissue and temporal-specific regulation to the individual globin genes and contains numerous DNase I hypersensitive sites, a hallmark of regulatory elements (Li et al. 1999). One of these sites (HS4) is able to block enhancers and prevent position effects, the latter of which is dependent on CTCF (Recillas-Targa et al. 2002). This site is also associated with the nuclear matrix, which is also dependent on CTCF (Yusufzai and Felsenfeld 2004). The CTCF binding site is also responsible for blocking of the spread of histone acetylation (Zhao and Dean 2004). Conserved CTCF sites flank the entire β -globin domain and may serve as delimiters of the domain from outside influence (Saitoh et al. 2000). Before the looping structures at the imprinted Igf2/H19 locus were discovered (Murrell et al. 2004), a similar looping structure was found for the LCR and the transcribed globin genes (Carter et al. 2002). The β -globin LCR has shown the way in the study of imprinting and other epigenetic phenomenon, and parallels continue to come to light. In a way, ICs can be thought of as LCRs that confer allele instead of tissue-specific regulation of distant genes.

From insulator to "imprintor"

The discovery that CTCF not only bound at the *Igf2/H19* DMR, but did so depending on methylation, gave important insights into CTCF function (Bell and Felsenfeld 2000; Hark et al. 2000). This became the first chromatin insulator controlled by DNA methylation, linking yet another epigenetic mark to CTCF other than HDAC recruitment. This link between DNA methylation and CTCF binding is especially relevant

to cancer, where genome-wide hypomethylation may cause spurious CTCF binding and cause dysregulation of many genes (Mukhopadhyay et al. 2004). In fact, mutations in cancers have been characterized in CTCF that change its binding specificity (Filippova et al. 2002). This discovery also identifies CTCF as a trans-acting factor for the regulation of imprinting, at least at *Igf2/H19*. It has been suggested, but not proven, that sites also exist at the PWS/AS IC (Ohta et al. 2001) and at the *Dlk1-Gtl2* locus (Takada et al. 2002). A role for CTCF has also been found for the regulation of X-inactivation (Chao et al. 2002).

The story of CTCF and imprinting takes an interesting turn with the cloning of a new paralog that shares the same ZF domains, but differs in C and N terminal domains (Loukinov et al. 2002). Named *BORIS* (brother of regulator of imprinting), its most striking characteristic is that it is expressed in a mutually exclusive manner to CTCF. While *BORIS* is expressed in spermatocytes, *CTCF* is expressed in spermatogonia, spermatids and all other somatic tissues. It is expected that BORIS has the same DNA binding spectrum as CTCF, but be able to carry out different biological activities at those sites. This and the fact that it is only in the testis during the erasure of DNA methylation in spermatogenesis, suggests that BORIS may have a very important role in imprinting, and more specifically, during the initial setup of imprints and differentially methylated domains. Other evidence indicates *BORIS* also has an important role in cancer (Klenova et al. 2002). While CTCF has so far not been observed in other imprinted domains, evidence suggests the future work on the *CTCF/BORIS* brothers will reveal their roles in many aspects of gene regulation, epigenetics, and perhaps a more general role in imprinting.

Objectives and Rationale

Much is known about the genetic function of the PWS/AS IC in determining the allelic identity for genes spread out over several Megabases (Nicholls and Knepper 2001). This involves many aspects of epigenetic control including DNA methylation, histone modifications and chromatin conformation. Regulation through an antisense mechanism involving a long paternal transcript originating from the IC region is an elegant explanation and has received much support (Runte et al. 2001). The story of imprinting in the PWS/AS cluster is far from complete however as the paternal expression of the transposed imprinted genes, *NDN*, *MAGEL2*, *MKRN3* and *Frat3* falls outside the scope of this antisense model. Therefore, I have sought to uncover the epigenetic mechanisms underlying imprinting of the transposed imprinted genes. My approach will seek to understand imprinting at different scales and how imprinted regulation intersects with tissue-specific regulation.

Focus will begin at the level of an individual imprinted gene. Previous work in our laboratory has already shown differential chromatin accessibility at *Ndn* (Hanel 2003). A fine scale analysis of the regulatory elements of *NDN* will further indicate accessibility of the two alleles. This will allow an examination of footprints for possible factors involved in allele specification as well as tissue-specific regulation in tissues that do and do not express *NDN*. Chapter 3 will follow my *in silico* analysis of the promoter, and collaborative work with Dr. Meredith Hanel on the *in vivo* footprinting assays in *NDN*.

With the advent of ChIP, covalent modifications of histones in *NDN* can also be studied, but can be done at a scale of the entire transcription unit as well as surrounding regions. This will allow examination of the chromatin context *NDN* is in on either allele,
and in different tissue types. It is conceivable that different histone codes specify imprinted versus developmental regulation. This study will also allow comparisons to histone modifications at the IC. It has been poorly characterized whether epigenetic characteristics at the IC are due to the imprinting function of the IC or transcription of the IC-associated transcription unit. Study of *NDN*, which is well isolated from the IC and is tissue-specific in expression, will help answer this question. Chapter 4 will reveal differences in the histone code correlated with allele-specific and tissue-specific regulation of *NDN*, using ChIP on human patient cells.

From the single representative transposed imprinted gene, it will be of great importance to evaluate if the other transposed imprinted genes are similar in their epigenetic regulation, as well as if these are conserved. If there are commonalities, perhaps mechanisms exist to co-regulate these genes. With the proposed role of CTCF in imprinting and function in long range control elements such as LCRs and insulators, it will be of value to study its role at the PWS/AS locus. This will aid in the understanding of locus-wide control of the transposed imprinted genes and give clues as to how their regulation fits with genes under control of the antisense transcript, as well as other imprinted genes. Chapter 5 will show evidence of an alternate mechanism of the regulation of the transposed imprinted genes that may be in a similar thread to that of the *lgf2/H19* locus.

The experiments presented here will also culminate in a more complete model of the regulation of the transposed imprinted genes, from allele-specific large scale domain regulation through CTCF, to local chromatin changes in histone modifications and factor binding reflective of tissue-specific differences. This will be discussed in Chapter 6.

Chapter 2 Materials and Methods

Parts of this chapter have appeared in:

- Hanel ML, Lau JC, Paradis I, Drouin R, Wevrick R (2005) Chromatin modification of the human imprinted NDN (necdin) gene detected by in vivo footprinting. J Cell Biochem 94(5):1046-57
- Lau JC, Hanel ML, Wevrick R (2004) Tissue-specific and imprinted epigenetic modifications of the human NDN gene. Nucleic Acids Res 32:3376-3382
- Lau JC, Wevrick R (Submitted) CTCF binds differentially methylated regions in the imprinted mouse Prader-Willi Syndrome locus.

Sequence analysis

Transcription factor binding prediction

Promoter sequences of *NDN/Ndn* from human, mouse and rat were compared using ClustalW (Thompson et al. 1994) at the European Bioinformatics Institute (http://www.ebi.ac.uk/clustalw/) and Pairwise BLAST (Tatusova and Madden 1999). Percent identity plots were performed with Microsoft Excel, using ClustalW data. Prediction of transcription factor binding sites was performed with Genomatrix MatInspector (Quandt et al. 1995). Positions -495 to +193 with respect to the start codon in human NDN, -553 to +222 in mouse *Ndn*, and -554 to +223 in rat Ndn were analyzed for sequence conservation and putative transcription factor binding sites.

CTCF binding cluster prediction

Genomic fragments from the February 2003 freeze of the annotated sequence from the UCSC Genome Bioinformatics Browser (http://genome.ucsc.edu/, chr7: 49276508-51906627, approximately 87 kb centromeric to *Snurf-Snrpn*, and 80 kb telomeric to *Frat3*) (Karolchik et al. 2003) were searched for the CTCF degenerate consensus CCGCNNGGNGGCAG (Chao et al. 2002) using pDRAW32 (http://www.acaclone.com/), reducing stringency to allow 3 additional mismatches to compensate for the broad binding abilities of CTCF. Regions containing four or more sites within 100 bp of each other were chosen for further analysis. Of these regions, only those with unique sequence amenable to PCR analysis were chosen. DNA from *Mus castaneus* and *Mus musculus* was sequenced to identify polymorphisms that were then used for allele-specific analysis. Direct sequencing of PCR products was performed using fluorescent dye terminator reactions and an ABI sequencer.

Tissues and cultured cell lines

The availability of human cell lines derived from PWS and AS patients with deletions of the 15q11-q13 region represented a unique reagent in which alleles can be studied in isolation. Control fibroblasts from the NIGMS Human Genetic Cell repository (GM00650), PWS fibroblasts (our laboratory number FB16, University of Miami Brain and Tissue Bank for Developmental Disorders #1889), and AS fibroblasts (FB17, 15q11-q13 deletion cell line, from Dr. A. Beaudet, Baylor College of Medicine) were grown in DMEM supplemented with 10% fetal bovine serum (FBS). Control lymphoblastoid cell lines (LCL) derived from primary blood lymphocytes (LCL10), PWS LCLs (LCL3, GM09024B, GM09133) and AS LCLs (LCL1/2, GM11515) were grown in RPMI supplemented with 15% FBS. Blood was collected from PWS and AS patients with fluorescence *in situ* hybridization-verified deletions and from control individuals, with informed consent.

Animal protocols were approved by the University of Alberta Health Sciences Animal Policy and Welfare Committee. Mouse strains *Mus musculus* (C57BL/6) and *Mus castaneus* (CAST) were obtained from the Jackson Laboratories, USA. We used polymorphisms between these two strains to distinguish parental alleles in F1 progeny. Animals were euthanized with euthanol (Sigma) and cervical dislocation. Tissues were processed for DNA, RNA, or ChIP immediately.

DNA extraction and PCR

Genomic DNA was extracted from tissue culture cells by proteinase K/SDS digestion, phenol/chloroform extraction and ethanol precipitation (Ausubel et al. 1993). Blood was collected in sodium EDTA tubes and erythrocytes lysed in two successive washes with 4 volumes of lysis buffer (1 mM EDTA, 10 mM KHCO₃, 155mM NH₄Cl, pH 7.4). After centrifugation, the cell pellet was washed with phosphate buffered saline (Wevrick and Francke 1996) and treated as above. DNA extraction from mouse tissues was performed with the DNeasy kit (#69506, Qiagen Inc).

PCR was performed in a PTC-100/200 thermocycler (MJ Research) with reagents from Invitrogen. Reactions are in a 20 μ l volume in thin walled PCR tubes. Primer sequences and conditions are in (Table 2-1).

Polymorphisms from regions of interest in human and mice strains (C57BL/6 and CAST) were identified by direct sequencing of PCR products and used to distinguish parental alleles.

RNA extraction and RT-PCR

RNA was extracted from tissues and cultured cells with Trizol reagent (Invitrogen) according to manufacturer protocols (Chomczynski and Sacchi 1987). Briefly, Trizol is an acid guanidinium thiocyanate and phenol solution that disrupts cellular membranes and proteins. Isolation of RNA is accomplished by chloroform extraction and isopropanol precipitation. Tissues required mild homogenization in a 1.5 ml eppendorf tube with a plastic disposable pestle (#199230001, ScienceWare). Total RNA was treated with DNase (Promega) and reverse transcribed (Invitrogen) into cDNA. This was used as template in

PCR reactions with gene specific-primers. Primers used in the following chapters are in (Table 2-1).

Nuclear extracts

Nuclear extracts were made by sequential lysis of cell and nuclear membranes in increasing salt (buffers NE1, 10 mM Hepes pH8, 1.5 mM MgCl2, 10 mM KCl, 1 mM DTT, and NE2, 20 mM Hepes pH8, 1.5 mM MgCl2, 25% glycerol, 420 mM NaCl, 0.2 mM EDTA, 1mM DTT, 0.5 mM PMSF) and dialyzed into NE3 (20 mM Hepes pH8, 20% glycerol, 0.1 M KCl, 0.2 mM EDTA, 0.5 mM DTT, 0.5 mM PMSF). Samples were flash frozen in a dry ice methanol bath and stored at -80°C.

Electromobility shift assay (EMSA)

EMSAs were done with the help of Gareth Cory in the laboratory of Dr. Alan Underhill. 6XHIS tagged, bacterially expressed paired-domains of Pax2 and Pax8 were kindly provided by Dr. Alan Underhill. ³²P labeled double-stranded oligonucleotides were incubated with protein or extract of interest in EMSA buffer (2.38X stock: 50 mM Tris 7.5, 250 mM KCl, 5 mM DTT, 10 mg/ml BSA, 50% glycerol), poly[dI:dC], and allowed to bind at RT for 30 minutes. Products were separated by polyacrylamide gel electrophoresis under non-denaturing conditions (6-10%). Oligonucleotide sequences are in (Table 2-1). Gels were dried and visualized by autoradiography (Biomax MR, Kodak).

Sodium bisulfite sequencing

Sodium bisulfite sequencing was used to identify specific methylated CpG residues in DNA. During this process, unmethylated cytosine residues in the template DNA are

converted to uracil, which is then converted to thymine during subsequent PCR. Bisulfite sequencing was performed as previously described (Hanel and Wevrick 2001), with minor modifications. Genomic DNA was treated with 0.3 M NaOH, 42°C/30 min, 95°C/3 min. The sample was then brought to 10X volume to 36.5% sodium bisulfite pH5 (wt/vol) and 1 mM hydroquinone and heated at 55°C overnight. The QIAEXII kit (Qiagen #20021) was used for DNA purification. PCR was performed as follows: (94°C/4 min, 58°C/2 min, and 72°C/2 min) repeated once, (94°C/30 sec, 58°C/30 sec, and 72°C/1 min) repeated 37 times, 72°C/10 min, hold at 4°C. Products were cloned into the pGEM–T vector (Promega Corp.). Ten to 30 individual clones of PCR products for each allele in each tissue were sequenced with dye terminators on an ABI sequencer. Primer sequences are in (Table 2-1).

Chromatin immunoprecipitation (ChIP)

The ChIP assay takes advantage of antibodies against specifically modified histones to enrich for DNA associated with such modifications (Figure 2-1) (Kuo and Allis 1999). While popularized by its utility in studying gene-specific histone modifications, it can also be used to study binding of non-histone chromatin proteins.

ChIP with human fibroblasts and lymphocytes

Chromatin immunoprecipitations were performed with reagents from Upstate Biotechnology (acetyl-histone H3 ChIP assay kit: #17-245, acetyl-histone H4 ChIP assay kit: #17-229, anti-dimethyl-histone H3 (Lys4): #07-030, anti-dimethyl-histone H3 (Lys9): #07-212, anti-dimethyl-histone H3 (K79): #07-366, ChIP assay kit: #17-295) and Abcam (anti-histone H3 (trimethyl K4): ab8580, anti-histone H3 (trimethyl K9): ab8898). The manufacturer's protocol was used with minor modifications. Crude lymphocyte preparations were made with Ficoll-Paque PLUS (Amersham Pharmacia Biotech) from 15 ml blood samples following the manufacturer's recommendations and expanded with phytohemagelutinin before being fixed as for other samples. LCL cultures and fibroblasts were fixed by addition of formaldehyde to a final concentration of 1% for 10 min at 37°C and cells collected. Fixed chromatin was sonicated with three 10 second pulses at one quarter maximum power with a 2 mm tip on a Fisher Scientific Sonic Dismembrator 60. Samples were pre-cleared with protein-A agarose beads prior to antibody addition. Mock control runs with no antibody were done and routinely gave no products from any primers used in this study. After reversal of the cross-links, DNA was extracted using commercially available binding columns (QIAquick PCR Purification Kit, Qiagen Inc.). The size of the resulting DNA fragments was determined by agarose gel electrophoresis. PCR was performed with one µl of template in each 20 µl reaction. SNRPN exon 1/IC primers were pair "A" as previously published (Saitoh and Wada 2000). PCRx (Invitrogen) was used as a PCR enhancer with a subset of primer pairs (see Supplementary Table). Quantification of ChIP experiments was done by densitometric analysis of ³²P end-labeled oligonucleotide probe hybridizations of slot-blotted PCR products detected on a Molecular Dynamics Typhoon and analysed with ImageQuant 5.2 quantitation software. Band intensities were corrected to background then normalized to GAPDH or chromosome 16 centromeric sequence (CEN16) bands before calculating a paternal versus maternal allele ratio. ChIP experiments were done in quadruplicate for fibroblasts, while limited availability of patient blood allowed only duplicate analysis. PCR amplifications and detection were performed

multiple times for each experiment. Primer and oligonucleotide probe sequences are in

(Table 2-1).



Figure 2-1. Chromatin immunoprecipitation (ChIP)

Covalent fixation between DNA and protein is caused by formaldehyde *in vivo*. Chromatin is then fragmented by sonication. Antibodies are used to enrich for nucleosomes carrying specific covalent modifications. Crosslinks caused by formaldehyde are reversed and DNA purified. Target sequences are detected by PCR amplification.

ChIP with mouse tissues

Chromatin Immunoprecipitation (ChIP) assays were performed according to manufacturer protocols (Upstate Biotech #17-295), with several modifications to the preparation of the starting material (nuclei isolation from tissues, Farnham lab, http://www.genomecenter.ucdavis.edu/farnham/farnham/protocols/tissues.html). Brain and liver were dissected from three week old F1 mice generated from a cross between a Mus castaneus male and a Mus musculus female. Tissues were chopped, homogenized with an A-pestle (loose) of a Dounce homogenizer in phosphate buffered saline (PBS) containing protease inhibitors (Mini Complete, Roche) with 1% formaldehyde, and fixed for 15 minutes at room temperature. Fixation was quenched with addition of glycine to 125 mM. Samples were further homogenized in cell lysis buffer (5 mM PIPES pH 8, 85 mM KCl, 0.5% NP40) in a Dounce homogenizer with the B-pestle (tight). Samples were strained through 3-4 layers of cheesecloth, collected by sedimentation (200 g, 5 min), and washed twice with cold PBS with protease inhibitors. Nuclei were lysed in ChIP kit lysis buffer, sonicated (twice, 10 sec each, power setting 2 on a Fisher Scientific Sonic Dismembrator 60), and specific proteins were immunoprecipitated according to a protocol from the manufacturer (Upstate). Antibodies were purchased from Upstate [anti-acetylated H3: #06-599, anti-dimethyl histone H3 (lysine 4): #07-030, anti-dimethyl histone H3 (lysine 9): #07-212, anti-dimethyl histone H3 (lysine 79): #07-366] and Abcam [anti-CTCF: ab10571, anti-trimethyl histone H3 (lysine 4): ab8580, anti-trimethyl histone H3 (lysine 9): ab8898]. Immunoprecipitated DNA was purified with the QIAEXII kit (Qiagen #20021). During the ChIP procedure, the chromatin is sonicated into fragments averaging 750 bp, so we

designed PCR primers that would effectively assay each CTCF binding site cluster but that did not necessarily include the predicted binding sites themselves. PCR was performed for each region and the resulting products were slot blotted on Hybond-N membrane (Amersham Biosciences RPN303 N). Enrichment of each allele was detected by hybridization with allele-specific oligonucleotides end-labeled with ³²P. Quantitation of signal intensity was performed using a Typhoon 8600 (Amersham Biosciences) and ImageQuant 5.2 software from the manufacturer. Allele-specific amplification and hybridization biases were controlled by similar quantitation of input DNA, defined to be 1% of starting material. We used PCR amplification of DNA from each parental strain on each blot to control for allele specificity of probes. A Student's T-test was performed on paired data points of both alleles within the same amplification reaction with the null hypothesis that each allele is equally amplified and detected for any given template. Primer and oligonucleotide sequences are in (Table 2-1).

Modification of measurement technique

A change in the methodologies for measurement and numerical analysis were employed in Chapter 5 versus Chapter 4. In data collected for Chapter 4, a large frame of the same size was drawn around bands and the internal intensity calculated by the ImageQuant 5.2 software, with the pixels under the frame itself taken as background (local average background correction, top left example, Figure 2-2). This was sensitive to fringing effects, where non-specific labeling of edges would occur, even in wells with no sample. For data in Chapter 5, tight frames were drawn around the band only, with no background correction (top right band). Background correction was found not to be necessary because

the background within a blot did not usually vary across the same blot. Shown below is an example of a blot for ChIP data in Chapter 5, with tight frames on each band.

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Wide border, LA correction



Narrow border, no correction





Figure 2-2. Change in measurement methods.

Example of modified measurement technique used in Chapter 5. Top left is an example of measurement used in Chapter 4 with a large frame and local average background correction. Top right is an example of measurement used in Chapter 5 with a narrow frame and no background correction. The top and middle rows show band examples described in text. Below is an example blot with narrow frames drawn.

A. CHAPTER 3 Oligonucleotides

EMSA oligonucleotides

Site	Forward Primer	F seq	Reverse primer	R seq
NDNPAX5	NDNPAX5-F	gcg cgg tcg cgc agg cgc agt gcc gcg tcc cg	NDNPAX5-R	gcg gga cgc ggc act gcg cct gcg cga ccg cg
PAX258	PAX258-F	gag ttg tga cgc act gaa gcg tga cga ctg tct	PAX258-R	gag aca gtg gtc acg ctt cag tgc gtc aca act
NDNCPBP	NDNCPBP-F	gtc ccg ccg ccg ccc cgc cct gcc cgt cgc tgc gg	NDNCPBP-R	gcc gca gcg acg ggc agg gcg ggg cgg cgg
NDNMOK	NDNMOK-F	gca gaa caa agt aag gat ctg agc gac cct aac ttt	NDNMOK-R	gaa agt tag ggt cgc tca gat cct tac ttt gtt ctg

RTPCR primers

I'll or pline								
Amplicon	Forward primer	Fseq	Reverse primer	R seq	Temp	MgCl	Cycles	Product size
PAX2	PAX2 RT-1F	cac aga cat acc ccg ttg tg	PAX2 RT-2R	tcg ttg tag gcc gtg tac tg	55	2.0	30	181
PAX5	PAX5 E5/6-F	gac gaa ggt att cag gag tc	PAX5 E10-R	cca agg gtc agt gac ggt c	55	2.0	30	585
PAX8	PAX8 RT-1F	gca acc att caa cct ccc ta	PAX8 RT-2R	ctg ctg ctg ctc tgt gag tc	55	2.0	30	240
CPBP	COPEB-1F	cac gag acc ggc tac ttc tc	COPEB-2R	ctg acc aaa act tcg cca at	55	2.0	30	359
EGR-1	EGRL-MMF	cag cag tcc cat tta ctc ag	EGRL-MMR	gac tgg tag ctg gta ttg	55	2.0	30	344
MAZR	ZNF278-1F	gcg ccy ata taa tgc tct tt	ZNF278-2R	tac cac ata gac cgc atg ga	55	2.0	30	390
hsMOK2	MOK2-1F	tee etg age tge aaa ett et	MOK2-2R	att gta ctg_gcc_atg_cct_tc	55	2.0	30	352

B. CHAPTER 4 Oligonucleotides

ChIP Primers

Amplicon	Forward primer	F seq	Reverse primer	R seq	Temp	MgCl	Cycles	Product size	PCRx
A	NEC122F	GGC CTA TTG CTA TGC CTG TC	NEC123R	CAC AGA GGC TGT CTC CCT TC	60	2.5	33-35	200	
В	NEC80F	TCC TCT CAC TGG TTC GCA TA	NEC81R	TGG GCT GAG AAG ATC TAG GG	55	2.0	33-35	219	
с	NEC87F	CCT GCC CTA GAT CTT CTC AGC	NEC34R	GGG GCC TCG GCT GCA AAG TTA GG	60	2.0	33-35	352	2X
D	NEC21F	GCG CAG ACA TGT CAG AAC AA	NEC69R	TTG ACC AGC ACG TAC CAC AT	50	3.0	33-35	343	3.5X
Е	NEC16F	ACG AGC TCA TGT GGT ACG TG	NEC17R	GAA GGT GGA GTG CTT CTT CC	50	2.5	33-35	376	2X
F	NEC20F	GCC CGA ATA CGA GTT CTT TT	NEC6R	CAC ACA TCA TCA GTC CCA TA	55	2.5	33-35	540	1
G	NEC120F	TGT GAG CAC TTG GCA CAC TT	NEC121R	GCG ATT TTT CCC ACC CTA TT	50	2.0	33-35	211	
н	NEC137F	GGC AGA AAA ACA ATG GAA GC	NEC138R	TTG TTT CTT TGT ACT ATT TTT CCT TT	C 60	2.5	33-35	117	1
SNRPN A	SNRPN AF	GAT GCT CAG GCG GGG ATG TGT GCG	SNRPN AR	GCT CCC CAG GCT GTC TCT TGA GAG	60	2.5	33-35	172	
GAPDH	GAPDH F	GCA TCA CCC GGA GAA ATC GG	GAPDH R	GTC ACG TGT CGC AGA GGA GC	60	2.5	33-35	268	
CEN16	CEN16 F	GTC TCT TTC TTG TTT TTA AGC TGG G	CEN16 R	TGA GCT CAT TGA GAC ATT TGG	55	2,5	33-35	207	

ChiP Oligonucleotide Probes

A	NEC176	cac	cac	caa	aag	ccc	ttt	ta		
в	NEC177	gag	aag	ggg	cca	gtt	taa	gg		
С	NEC178	gca	aag	tta	ggg	tcg	ctc	ag		
D	NEC68	GAG	CGA	CCC	TAA	CTT	TGC	AG		
E	NEC14	ATG	CTC	CTG	CAC	CAC	TTC	тт		
F	NEC2	ATT	TGC	ATC	TTG	GTG	ATT	т		
G	NEC8	GGT	GGG	GTT	GTA	TAT	GTG	тт		
н	NEC179	ttg	aaa	caa	gtt	ttt	gct	tcc	a	
SNRPN A	SNRPN A O	tga	cgc	atc	tgt	ctg	agg	ag		
GAPDH	end-labeled	PCR pro	duct	:						
CEN16	end-labeled	PCR pro	duct	2						

Table 2-1. Tables of oligonucleotides.Oligos used in various experiments in (A) Chapter 3, (B)Chapter 4, (C) Chapter 5.

C. CHAPTER 5 Oligonucleotides

ChIP Primers		
		size MgCl temp cycle
Frat3	cca ttc agt ggg tgt cag aa	222 2.5 60 33-35X
	aga atg ggc atc tga gac aa	222 2.5 60 33-35X
Mkrn3	gac agt gtc cct gcc aaa g	205 2.5 60 33-35X
	ggc aaa gtc cag gct tct ac	205 2.5 60 33-35X
Ndn	cca tca tct agt tct gtg cca	284 1.5 60 33-35X
	ctt cgg atc aga gca gga c	284 1.5 60 33-35X
ict-A	aag tgg ttg gcc ttg tct gt	142 2.5 60 33-35X
	ttc aga atg aca gtt cac att gc	142 2.5 60 33-35X
iCT-B	agg aac act tgt ggc ttg aga	217 2.5 60 33-35X
	gaa aca cac tgc agc agc tc	217 2.5 60 33-35X
Snurf/Snrpn/IC	caa cag agc tcc tgc atc ct	226 2.5 60 33-35X
	gcc tct gga ctc ctg gaa g	226 2.5 60 33-35X
Dadl exl'	ggg cag cag tac tcc acc aa	159 2.5 60 33-35X
	cgt agg atg cag gga ttt tct tta	159 2.5 60 33-35X
Dadl ex6	ccc aca gat tga aca cag gaa at	179 2.5 60 33-35X
	gag gga tog atg tet get gtt t	179 2.5 60 33-35X

RT-PCR Primers (Frat3 and Mkm3 requires PCRx)

								size		MgCl	temp	cycle	PCRx
Frat3	gac	gtg	gac	cgg	ctc	atc			509	2.5	55	30-35X	2X
	agg	ttt	ccc	gaa	aga	agg	ag		509	2.5	55	30-35X	2X
Mkrn3	caa	gcc	ttg	cag	cag	gtg			298	2.5	60	30-35X	1X
	att	tct	сса	tgg	3 33	tat	gc		298	2.5	60	30-35X	1X
Ndn	gta	tcc	caa	atc	cac	agt	gc		356	2	55	30-35X	
	Ctt	cct	gtg	cca	gtt	gaa	gt		356	2	55	30-35X	
ict-a	cag	cag	agg	act	tcc	tgg	tc		150	1.5	. 60	30-35X	
	tgg	ggt	gtt	aat	tcc	acg	tt		150	1.5	60	30-35X	
Snurf/Snrpn/IC	CCC	GAG	TAT	TAA	GGA	TCT	TG		142	1.5	55	30-35X	
	TGA	AGA	TTC	TCC	CAT	CTT	GC		142	1.5	55	30-35X	
Gapdh	GCC	ATC	AAC	AAC	ccc	TTC	AT		315	2	60	30-35X	
	TTC	ACA	CCC	ATC	ACA	AAC	АТ		315	2	60	30-35X	

Allele-specific oligonucleotide probes

									allele
Frat3	tct	taa	agt	cag	att	aca	g		Cast
	tct	taa	agt	aag	att	aca	g		C57
Mkrn3	cct	tcg	acg	ctt	gta				Cast
	cct	tca	atg	ctt	gta				C57
Ndn	tgc	ttc	gct	cct	ttc				C57
	tgc	ttc	ggt	cct	ttc				Cast
iCT-A	cgg	cct	cca	tct	ag				Cast
	cgg	ctt	сса	tct	ag				C57
iCT-B	ccc	tgc	cta	gcc	ct				C57
	ccc	tgt	cta	gcc	ct				Cast
Snurf/Snrpn/IC	aac	ttc	tac	cca	cac	cc			Cast
	aac	ttc	tat	cca	cac	cc			C57
Dadl exl'	ctc	ggc	gtg	ttt	att	с			
Dadl ex6	cac	aca	cac	aca	σaσ	aaa			

Bisulfite Sequencing Primers

Frat3	required	semi-n	este	d PC	R, f	irst	rou	nd w	ith	F-fl	ank/R	and	second	_round	with	F-nest/R
													:	size		
Frat3		gag	ttt	ttt	tgg	tgg	taa	tga	tta	ga		F-f	lank	746		
		ttt	taa	tta	aga	atg	aga	agt	tta	ggt	tt	F-r	lest	676		
		aaa	tac	tta	ctt	tac	cca	tcc	cc			R				
Mkrn3		ggg	taa	ttg	aat	ttg	ttt	ttg	gat	а				238		
		taa	aaa	tat	aca	cac	cta	tcc	cca	с				238		
iCT-A		aaa	aaa	act	ccc	aca	aca	caa	taa	с				312		
		ttt	tta	ggt	tta	ątt	aaa	ttt	ttt	aga	tta		-	312		

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Chapter 3 Analysis of the NDN Promoter

Parts of this chapter have appeared in:

Hanel ML, Lau JC, Paradis I, Drouin R, Wevrick R (2005) Chromatin modification of the human imprinted NDN (necdin) gene detected by in vivo footprinting. J Cell Biochem 94(5):1046-57

Introduction

Gene regulation involves complex networks of *cis*-acting genetic elements and *trans*-acting protein factors. The most tangible of these *cis*-acting elements is the promoter of a gene. Binding and activation of RNA polymerase and associated transcriptional machinery is the ultimate end result of a concert of regulatory events that includes the availability of tissue-specific transcription factors as well as accessibility of the locus in terms of its chromatin state. For imprinted genes that are also transcribed in specific spatial-temporal patterns, such as *NDN*, *MAGEL2*, and *MKRN3*, an extra layer of regulation must intersect at the promoter to control correct allele-specific expression. Detailed examination of the promoter of these genes may give clues as to chromatin accessibility as well as the *trans*-acting factors involved.

NDN is expressed at varying levels in many human tissues but is highest in brain, heart, muscle and fibroblasts. However, its expression is not ubiquitous and *NDN* is not expressed at appreciable levels in liver, kidney and blood leukocytes (Jay et al. 1997; MacDonald and Wevrick 1997). Mouse *Ndn* is also tissue-specific, and is expressed in a pattern that partially overlaps the human *NDN* expression profile (Uetsuki et al. 1996). In particular, it is more neurally restricted. Since *NDN* was localized to the PWS critical region, it became an excellent candidate gene for phenotypes in the disease because it had been implicated in terminal differentiation of neurons (Uetsuki et al. 1996; MacDonald and Wevrick 1997), respiration (Ren et al. 2003), and axon outgrowth (Lee et al. 2005). While the expression profiles of human and mouse *NDN/Ndn* are not identical, the similarities suggest overlap in aspects of tissue-specific regulation. Both orthologues are imprinted in

all expressed tissues examined, and therefore may share mechanisms of allele-specific expression.

Previous analysis of the human and mouse *NDN/Ndn* promoters involved reporter construct transfections into cultured cells (Uetsuki et al. 1996; Nakada et al. 1998). This group defined the minimal promoter in both species and found that the mouse promoter conferred expression of a reporter gene in post-mitotic neurons during *in vitro* differentiation of P19 cells. It was also found that the human promoter was sensitive to DNA methylation, in that methylation of seven *HhaI* sites greatly reduced the ability of this promoter to activate the reporter. In a comparison of the DNA sequences of the two promoters, a site resembling the binding site for the transcription factor SP1 was found in both sequences in corresponding locations (Nakada et al. 1998). In this chapter, I further analyzed the DNA sequence of the promoters and characterized other potentially important sites where *trans*-acting factors may bind to mediate tissue or allele-specific transcription.

Footprinted sequences in the NDN promoter

In parallel to this work, Dr. Meredith Hanel of our laboratory had undertaken *in vivo* footprinting studies of the *NDN* promoter (Hanel et al. 2005). Briefly, deletion patient cell lines from PWS and AS patients were used in DNase I, DMS and UVC footprinting assays. The use of deletion patient cells allowed unequivocal differentiation of alleles, with the maternal allele remaining in PWS, and paternal allele remaining in AS. Fibroblasts and lymphocytes were used to represent *NDN* expressing and non-expressing tissues, respectively. These studies enabled a fine scale ascertainment of chromatin access on either allele in expressing and non-expressing contexts. I used the data generated by Dr. Hanel to

examine promoter occupancy, in comparison with bioinformatics analysis of the promoter, to identify candidate factors regulating allele-specific and tissue-specific transcription of *NDN*.

One of the footprints Dr. Hanel found was a striking paternal allele-specific DNase I footprint on both strands, located in the promoter between -136 and -166 bp upstream of the start codon (Hanel et al. 2005). This was only found in fibroblasts, and therefore may correspond to transcription-related factors used by the expressing allele. The occupancy of this site was supported by the presence of UVC and DMS footprints as well. This indicates that a protein or protein complex is bound to the active allele, covering both strands over a region of about 30 bp with high GC content. This footprint will be referred to as the transcription-related footprint.

Conservation of sequence features between these promoters despite the 75 million years between human and mouse (Waterston et al. 2002) suggests a functional role and will lead to a better understanding of the regulation of *NDN/Ndn* (Pennacchio and Rubin 2001). To this end, this chapter will describe comparative sequence analysis of the *NDN/Ndn* promoter of human, mouse, and rat. These phylogenetic footprints were correlated with *in vivo* footprints performed by Dr. Hanel. Overlaps between phylogenetic and *in vivo* footprints are of special interest, and were investigated in terms of sequence motifs previously characterized to be binding sites of transcription factors. Potential binding of a candidate factor was studied using *in vitro* binding assays. The data suggest that the differential footprinted patterns observed between expressed and non-expressed alleles may be due in part to accessibility of the chromatin at the promoter region and not solely to the availability of *trans*-acting factors.

Results

Phylogenetic analysis of the NDN/Ndn promoter

To study the sequence conservation between human, mouse and rat NDN/Ndn, which may be indicative of conserved functional elements, the upstream regions from the three species were compared using ClustalW as implemented by the European Bioinformatics Institute (Figure 3-1) (Thompson et al. 1994). By visual inspection, it is clear that the degree of similarity between mouse and rat sequences is very high, as expected for the closely related species. To further quantitate the sequence similarity between the three sequences, a percentage identity plot was produced from the ClustalW data, using a 50 bp window (Figure 3-2 A). As can be seen from the plot of the alignment between all three sequences, the similarity drops dramatically a short distance upstream of the transcription start. Since the mouse and rat sequences retain more extensive sequence similarity, the change in similarity among all three sequences is attributable to differences between human and rodent sequences. Closer inspection of the sequence alignment (Figure 3-1) shows a well defined point at which the level of similarity changes at position -133. In fact, overall sequence similarity 5' to this is 40.5%, while it is 69.1% 3' of this position, perhaps suggesting an evolutionary rearrangement event between human and rodents. It is notable here that most of the previously defined minimal promoters in human and mouse are in the less conserved region (Figure 3-2 A).

			-439
Human	TTCCTCTCACTGGTTCGCATAAAGCTCATGTTTACAAAGCCGCCCAGACCTT-TCTC	56	-433
Mouse	CTCATCATCATCATCATAAGGTACAGCTTTCCAAAAGAAAAAAGAAAAAAAAGTCATTTC	105	-448
Rat	CACATCATCATCATCATCATCATCATCAGGTATGACTTTCCAAGAAAAGGTTATTTT	120	-435
3W	** ** ** ** ** * * * * * * *		
HM	** ** *** *** *** * * * * * * * *	-	
HR	*** *** * * * * ** * * * * * * * * * * *	-	
MR	* ********** * ** * * * * * * ** **** ** ****	-	
•			
•	· · · · · · · · · · · · · · · · · · ·		
Human	TGGGACTCTCATATTTAACTTAATTCTGGATATACCCAGGTAAGCGTTTCCCAAGAA	113	-382
Mouse	TGTTTCTCTTATTCTTTGTAGAAAAACCAAAATCAAGAATTAAGTCTTTCTCCAGGA	163	-390
Rat	TGTCTCTCCTATTCTTTGTGGAAAACTCAAATTCAAGAATTAGTCTTCTTCCTCCAGGA	180	-375
3W	•• ••• •• •• • • • • • • • • • • • • • •		
HM		-	
HR	· · · · · · · · · · · · · · · · · · ·	-	
MR	*** **** *********** ***** **** *******	-	
•			
Human	ACTTGACCCCAACATCCCAAAAACTTAAGGTATCTTTCCCTTAAACTGGCCCCT	167	-328
Mouse	CCTTCACATTTAATTTGATTTTGCACAAACTCAGTGTGTCCGAAC-TTAACTTCACCATC	222	-331
Rat	CCTTCRCACTTARCTCRGTGTGTGGGAAC-TTRACTTCRCCATC	223	-332
3W	*** ** *		
HM	*** ** * * ** ** ** ** * **** * ***	-	
HR	· ·· · · · · · · · · · · · · · · · · ·	-	
MR	******** ****	-	
Human	TCTCCRGTRCGCATCCATCTCACTTCTCTCC-TGCCCTAGATCTTCTCAG-CCCR	220	-275
Mouse	TCAGCTACATCTTCTCCCTTCAACTTCTTTCTTTCCCTACCATCAT	282	-271
Rat	TCAGCTATATCTTTCTCCTTCAACTTGTTTCTTTCCCCACTATCACCCATCTCTCTC	283	-272
34	** * * * * * * * * *** *** * * * * * * *		
HM	** * * * * * ** ** **** ** **** **	-	
HR .	** * ** ** *** * * **** * ****	_	
MP	******* *******************************	_	
nn i		-	
•			
•	ŧ		
Human	AACAGGAAACCCCGGGATCGCTCTCCCAGCAGGTGAAGCCTCGCCATGGACCCTCCCC	278	-217
Mouse	TACAGGAGACCAGGAAATCTTTTACATAAGCCTAGTGGTACCCTCCCT	342	-211
Rat	TACAGGAGACCAGGAGATCCTTTTCATAATACTAGTGGTACCCTCCCT	343	-212
3W			
HM	****** *** * *** * * * * ** **	-	
HR	****** *** * **** * ** * *** *** *** ** ****	-	
MR	***************	-	
Ruman	GTEGGGGGECOGEGETGEGEGEGEGEGEGEGEGEGEGEGEGEGEGEGEGE	336	-159
Mouse	GTTGGGCTTTGGCTGGTTGGCTGGCTACCACCCTTCTGGCTTCCCAACACC	402	-151
Pat	GTTGGGCTTTGCTTGCTTGCTGCTGCCGTGCGCCTTCTGCCTTCCCGCCG	403	-152
714 L		403	-132
3W			
117		-	
nk ND		-	
MR		-	
•			
•			~~
Human	AGGCGCAGTGCCGCGTCCCGCCGCCCGCCCCGCCCGCCGCGCGCAAGGCGCCGCG		
		396	- 55
Mouse	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTGCTGCGGAAGGCGCAGTG	396 454	-99
Mouse Rat	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTGCTGCGGAAGGCGCAGTG ATGCGCAATATAGCGTCTGCCCCGCCCGGCCCGGC	396 454 456	-99 -99
Mouse Rat 3W	ATGCGCRATATCGCATCAGCCCCGCCCGCCGCTGCTGCGGAAGGCGCAGTG ATGCGCAATATAGCGTCTGCCCCGCCCGGCCCGG	396 454 456	-99 -99
Mouse Rat 3W HM	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCGCAATG ATGCGCAATATAGCGTCTGCCCCGCCCGCCCGCCGCTGCTGCGGAAGGCGCAATG	396 454 456 -	-99 -99
Mouse Rat 3W HM HR	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTGCTGCGGAAGGCGCATG ATGCGCAATATAGCGTCTGCCCCGCCCGGCCCGGCC	396 454 456 - -	-99 -99
Mouse Rat 3W HM HR MR	ATGCGCARTATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCGAGTG ATGCGCARTATACGCTCTGCCCCGCCGGCCCGCCCGTGCTGCGGAAGGCCCAGTG	396 454 456 - -	-99 -99
Mouse Rat 3W HM HR MR -	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCGCAATG ATGCGCAATATAGCGTCTGCCCCGCCCGGCCGGCCGCTGCTGCGGAAGGCGCAATG	396 454 456 - -	-99 -99
Mouse Rat 3W HM HR MR	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCGCAGTG ATGCGCAATATACGCATCTGCCCCGCCGGCCCGCCTGCTGCGGAAGGCCCCAGTG 	396 454 456 - -	-99 -99
Mouse Rat 3W HM HR MR • • •	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCGATG ATGCGCAATATACGCTCTGCCCCGCCGGCCGCCGCTGCTGCGGAAGGCCGATG 	396 454 456 - - - 456	-99 -99
Mouse Rat 3W HM HR MR Human Mouse	AT6C5CAATATC5CATCA5CCCC5CCCC5-CCC5CT6CT6C5GAA565C5CA5T6 AT6C5CAATATA5C5CTC5CCCC5CC5C6CC5CCT6CT65GAA665C5CA5T6 	396 454 456 - - - 456 514	-99 -99 -99
Mouse Rat 3W HM HR MR Human Mouse Rat	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCCAATG ATGCGCAATATCACGTCGCCCCGCCGGCCCGCTGCTGCGGAAGGCCCAATG 	396 454 456 - - 456 514 516	-39 -39 -39
Mouse Rat 3W HM HR HR - Mouse Rat 3W	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTGCTGCGGAAGGCGCAATG ATGCGCAATATACGCTCTGCCCCGCCGGCCGCCGCTGCTGCGGAAGGCGCAATG 	396 454 456 - - - 456 514 516	-39 -39 -39
Mouse Rat 3W HM HR MR MR Human Mouse Rat 3W	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGAAGGCCCCATG ATGCGCAATATACGCATCAGCCCCGCCGGCCCGCTCCTGCGAAGGCCCCATG 	396 454 456 - - - 456 514 516 -	-39 -99 -39 -39 -39
Mouse Rat 3W HM HR MR Muse Rat 3W HR HR	ATGCGCAATATCGCATCAGCCCCGCCCGCCGCCGCCGCCGCCGCGCGGCGGC	396 454 456 - - 456 514 516 -	-39 -39 -39 -39 -39
Mouse Rat 3W HM HR MR MR Human Mouse Rat 3W HM HR HR HR	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGAAGGCCCAATG ATGCGCAATATACGCTCTGCCCCGCCGGCCCGCTGCTGGGAAGGCCCAATG 	396 454 456 - - 456 514 516 - -	-99 -99 -99 -39 -39 -39
Mouse Rat 3W HM HR MR MR Nouse Rat 3W HM KR	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCCATG ATGCGCAATATACGCTCTGCCCCGCCGGCCGGCCGGTGCTGCGGAAGGCCCATG 	396 454 456 - - 456 514 516 - -	-99 -99 -39 -39 -39
Mouse Rat. 3W HM HR MR MR Human Human Mouse Rat. 3W HM HR HR HR	AT6C6CAATATC6CATCA6CCCC6CCCCG-CCC6CT6CT6C6GAA66CCCAATG ATCCCCAATATAC6CTCT6CCCC6CCC6CCCC6C	396 454 456 - - - 456 514 516 - -	-39 -99 -39 -39 -39
Mouse Rat 3W HM HR MR MR Human Mouse Rat 3W HM HR KR KR KR KR KR KR KR KR KR KR KR KR KR	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTCCTGCGAAGGCCCCATG ATCCGCAATATCGCATCAGCCCCGCCCGCCCGCTCCTGCGAAGGCCCATG 	396 454 456 - - - 456 514 516 - - - 516	-39 -99 -99 -39 -39 -39 -39
Mouse Rat 3W HM HR MR • • Human MR HR HR HR HR HR Human Mouse	AT605CAATAT06CAT0A6C0000005-C005CT0CT0506AA565C5CAT0 AT000000000000000000000000000000000	396 454 456 456 514 516 516 574	-39 -99 -39 -39 -39 -39 -39 +21 +21
Mouse Rat 3W HM HR MR MR Human Mouse Rat 3W HR MR HUMAN MR Rat	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTGCTGCGGAAGGCCCGATG ATGCGCAATATCGCATCAGCCCCGCCGCGCGCCGTGCTGCGGAAGGCCCAATG CCCCGCACGCGCCTTCTTCTCCCAGGAATCCGCGGAGGGGCGCAGGGCCGAAGGCCC CTGCGAAGGCGCCGCTTCCTCCTGCGAGGACTCCGCGGAGGAGGCCCAGGGCCCGAGGAGGCC CTGGACGGAGGGCCCTGCCCTGCCCAGGACGCGCGAGACATGTCAGAAGAAGTAAGGAC CAGGCGCGAGGGCCCTGCCCTGCCCAGGAGGCCCAGACATGTCAGAAGAAGTAAGGAC CAGGCGCGAGGGCCCTGCCCTGCCCAGGAGGCCCAGACATGTCAGAACAAAGTAAGGAC CAGGCGCGAGGGCCCTGCCCTGCCCTGCCGAGGAGCCGAGACATGTCAGAACAAAGTAAGGAC	396 454 456 - - - 456 514 516 - - 516 574 575	-39 -99 -99 -39 -39 -39 -39 +21 +21 +21
Mouse Rat 3W HM HR MR MR MR Rat 3W HM MR HR MR HR MR HR MR MR MR SW SW SW	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCCAATG ATGCGCAATATCGCATCAGCCCCGCCGGCCGGCCGTGCTGCGGAAGGCCCAATG 	396 454 456 - - - 456 514 516 - - 576 574 577	-39 -99 -99 -39 -39 -39 +21 +21 +21
Mouse Rat. 3W HM HR MR MR HIMAN Mouse Rat 3W HM HR HM HIR NR HUMAN HUMAN HUMAN HUMAN HUMAN HUMAN HIM	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCCAATG ATGCGCAATATCACGTCGCCCCGCCGGCCGCCGTGCTGCGGAAGGCCCAATG 	396 454 456 456 514 516 576 574 575 -	-39 -99 -99 -39 -39 -39 +21 +21 +21
Mouse Rat 3W HM HR MR MR HR Auman Mouse Rat 3W HM HR MR SW HUMAN MOUSE Rat 3W HUMAN HR	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCCAATG ATGCGCAATATCGCATCAGCCCCGCCGCCCGCCCGTCCTGCGGAAGGCCCAATG 	396 454 456 516 5516 516 574 575 	-39 -99 -99 -39 -39 -39 +21 +21 +21
Mouse Rat 3W HM HM HR MR HR Human Mouse Rat 3W HM Human Mouse Rat 3W HIM HR MR	AT6C6CAATATC6CATCA	396 454 456 - - - - - 516 574 575 - - - -	-99 -99 -99 -39 -39 -39 +21 +21 +21
Mouse Rat 3W HM HR MR MR Human Mouse Rat 3W HM HR MR KR SW HUMAN MR HUMAN MR SW HUMAN	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTGCTGCGGAGGCCCAATG ATGCGCAATATACGCATCAGCCCCGCCGCCGCCGCTGCTGCGGAGGCCCAATG 	396 454 456 456 514 516 	-99 -99 -99 -39 -39 -39 +21 +21 +21
Mouse Rat 3W HM HR MR MR HR Human Mouse Rat 3W HM Human Mouse Rat 3W HIR HIR HIR HIR SA SA HI HIR HIR	AT6C6CAATATC6CATCA	396 454 456 - - - 516 574 575 - - -	-99 -99 -99 -39 -39 -39 +21 +21 +21
Mouse Rat 3W HM HR MR MR M Human Mouse Rat 3W HM HR MR MR HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTCGTGCGGAAGGCCCAATG ATGCGCAATATCGCATCAGCCCCGCCCGCCGCCGTCGTGCGGAAGGCCCAATG ATGCGCAATATCGCGTCTTTCTCCCAGGAAGCCGCGGAGGGGCCGCAGGGCCGGAGGGGCCGAAGGCCCAATG CGCAGCAACGCGCACTTCCTCTCCGCAGGCGCCGCAGGGGGCCGCAGGGGCCCGAGAGGGCCCGAGGGCCCCGAGAGGGCCCGGAGGA	396 454 456 556 5514 5516 	-99 -99 -39 -39 -39 -39 +21 +21 +21
Mouse Rat 3W HM HR MR Human HM HR HR Human HM Mouse Rat 3W HM HR HT Human HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCAGCCCCCCCCG-CCCGCTGCTGCGGAAGGCCCAATG ATGCGCAATATCGCATCAGCCCCCGCCGCCCGCTGCTGCGGAAGGCCCAATG ATGCGCAACAGCGCATCAGCCCCGCGCGCCCGCTGCTGCGGAAGGCCCAATG Image: Comparison of the second of the s	396 454 456 	-99 -99 -99 -39 -39 -39 +21 +21 +21 +21 +21
Mouse Rat 3W HM HR MR MR HR Human Mouse Rat 3W HM HR KR KR HUMAN MOUSE Rat 3W HM HR KR HUMAN HR Rat 3W HM HR Rat 3W HM HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCCAATG ATGCGCAATATCACGTCGCCCCGCCGCCGCCGCTGCTGCGGAAGGCCCAATG 	396 454 456 456 514 516 516 574 5775 576 637	-99 -99 -99 -39 -39 -39 +21 +21 +21 +21 +81 +81
Mouse Rat 3W HM HR MR MR Human Mouse Rat 3W HM MR HR MR Human HR MR HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTCGTGCGGAAGGCCCCATG ATGCGCAATATCGCATCAGCCCCGCCCGCCGCCGTCGTGCGGAAGGCCCAATG CCCGCAGCAATATCGCGTCT	396 454 456 - - 456 514 516 - - 576 574 575 - - 576 6334 635	-99 -99 -99 -39 -39 -39 +21 +21 +21 +21 +81 +81 +81
Mouse Rat. 3W HM HM HR MR Human Mouse Rat. 3W HM HR Rat. 3W HUMAN HR HR HR HR HR HR HIMAN HR HIMAN	ATGCGCARTATCGCATCA	396 454 456 - - 456 514 516 - - 576 575 - - 576 635 -	-39 -99 -99 -39 -39 -39 +21 +21 +21 +21 +81 +81
Mouse Rat 3W HM HR MR MR Mouse Rat 3W HM HR MR Human Mouse Rat 3W HM KR Kat 3W HM KR MR	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTGCTGCGGAAGGCCCAATG ATGCGCAATATCGCATCAGCCCCGCCGCCGCCGTGCTGCGGAAGGCCCAATG ATGCGCAACGCGCACTTCT-CTCCCAGGAATCCGCGGAGGGGCGCAGGGCCGAAGGCCCAAGGACCC CTCAGTAAAGCGCACTTCCTCTCCGCTGGCTCCCACCGAGGAGGCGCCAGGGCCCGAAGAGCCC CTCAGTAAAGCGCACTTCCTCTGCTGCCGCAGCGCGCAGGACATGTCAGGACGAAGGAC CTGGACGCAGAGGCCCTGCCCTTGCCAGGCGCGCGAGACATGTCAGGACAAAGTAAGGAT CAGGCGCCGCCTGCCCTTGCCCAGGCGCGCGAGACATGTCAGGACAAAGTAAGGAT CAGGCGCCGCGCCTGCCCTGCCCTGCCCAGGCGCCGCAGACATGTCAGGACAAAGTAAGGAT CAGGCGCCGCGCCTGCCCTGCCCTGCCAGGCGCCGCAGACATGTCAGGACAAAGTAAGGAC CAGGCGCCGCGCCTGCCCTGCCCTGCCCAGCCGCGCAGACATGTCAGGACAAAGTAAGGAT CAGGCGCCCGCCTGCCCTGCCCTGCCCAGCCGCCGCAGACATGTCAGGACAAAGTAAGGAC CCTGAGCGCCGCGCCTGCCCTGCCCTGCCCCGCCGCAGACATGTCAGGACAAAGTAAGGAC CCTGAGCGACCCTAACTTTGCAGCCGAGGCCCCCAACTCCGAGGTGCACGACACCGATGCC CTGAGCGACCCTAACTTTGCAGCCGCAGGCCCCCGACCTGGGGTGCAGCGACGCCCTGCC CTGAGCGACCCTAACTTTGCAGCCGCAGGCCCCCGCACCTGGGGTGCAGCGACGCCCTGCC CTGAGCGACCCTAACTTTGCCGCCTAGGCCCCGACCTGTGGGTGCCCCCACCTGCGCCTCCCCCGCCCTGCCCTGCCCTCGCGCCCGCC	396 454 456 - - 456 514 516 - - 516 574 575 - - 576 635 - -	-39 -99 -39 -39 -39 +21 +21 +21 +21 +21 +81
Mouse Rat JW HM HR MR MR Mouse Rat JW HM HR HR HR Mouse Rat JW HUMAN MR K HUMAN HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCSCARTATCSCATCA	396 454 456 - - 456 514 516 - 576 634 575 - 576 635 -	-39 -99 -99 -39 -39 -39 -39 +21 +21 +21 +81 +81
Mouse Rat 3W HM HR MR MR Mouse Rat 3W HM HR Mouse Rat 3W HM HR NR S S HM HR MOUSE Rat 3W HM HR MOUSE Rat 3W HM HR MOUSE Rat 3W	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCCAATG ATGCGCAATATACGCTTGCCCCGCCGCCGCCGTGCTGCGGAAGGCCCAATG CCCCGCACGCACGCCGTCTTCCCCGGGAACGAGGCCCGAAGGCCCGAAGGCCCCAAGGCCCCAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGAAGGCCCCGACGA	396 454 456 - - - 456 516 516 - - 516 574 575 - - 576 635 - -	-39 -99 -99 -39 -39 -39 +21 +21 +21 +81 +81
Mouse Rat JW HM HR MR · · Human Mouse Rat JW HM HR Human Mouse Rat JW Human Human Mouse Rat JW HUM HR HR HI HR HI HI HI HI HI HI HI HI HI HI HI HI HI	ATGCGCAATATCGCATCAGCCCCCCCCG-CCCCGCTCGTCGCGAAGGCCCCAGTG ATGCCCUARTATACCTTCGCCCCGCCCCGCCCGTCGTCGCGAAGGCCCCAGTG 	396 456 	-39 -99 -39 -39 -39 -39 +21 +21 +21 +81
Mouse Rat 3W HM HR MR MR Mouse Rat 3W HM HR KR KR Mouse Rat 3W HM HR MR	ATGCGCAATATCGCATCAGCCCCGCCCCG-CCCGCTGCTGCGGAAGGCCCAATG ATGCGCAATATAGCGTCGCCCCGCCGGCCGCCGTGCTGCGGAAGGCCCAATG 	396 454 456 	-39 -99 -99 -39 -39 -39 +21 +21 +21 +21 +81 +81
Mouse Rat JW HM HR HR MR Rat JW HM HR HR HR HI HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCSCAATATCGCATCAGCCCCCCCCG-CCCCGCCCCCCCCCC	3964 456 	-39 -99 -99 -39 -39 -39 -39 +21 +21 +21 +81 +81
Mouse Rat 3W HM HR HR MR HR HUMAN Mouse Rat 3W HM HR HC HUMAN MR H HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCARTATCGCATCA	3964 456 $ 4565$ $ 456514$ 5166 $ 57745$ $ 56345$ $ 576435$ $ 5694$	-39 -99 -99 -39 -39 -39 +21 +21 +21 +21 +81 +81 +81 +129 +141
Mouse Rat 3W HM HR MR MR Mouse Rat 3W HM MR HR HUMAN HR HUMAN HR HUMAN HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTCCTGCGGAAGGCCCAATG ATCCGCAATATCACGTC	3964 456 $ 4566$ 514 $ 5164$ 5755 $ 56345$ $ 5744$ $ 5744$ $ 5744$ $ -$	-39 -99 -99 -39 -39 -39 +21 +21 +21 +21 +81 +81 +81 +129 +141 +141
Mouse Rat SW HM HR HR MR · · Human Mouse Rat SW HM HR Rat SW HM HR Rat SW HM HR HR Rat SW HM HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCA	3964 456 $$ 4566 5114 $$ 5166 5775 $$ 576435 $$ 576435 $$ 62945	-39 -99 -99 -39 -39 -39 -39 +21 +21 +81 +81 +81 +81 +129 +141
Mouse Rat JW HM HR HR MR MC Ka Rat JW HM HR MC SM HM HR MC SM HM HR MC SM HM HR MC SM HM HR MC SM HM HR MC SW HM HR MC SW SW HM HC SW SW HM HC SW SW SW SW SW SW SW SW SW SW SW SW SW	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTCCTGCGGAAGGCCCAATG ATCCCGAATATACCGTTGCCCCGCCGCGCCGTCCTGCGGAAGGCCCAATG CCCGCACCAACCGCGATTTCTTCTCCCAGGAATCCGCGGAGGGCGCAGGGCCGAAGGCCCAAGGCT CTCAGTAAAGCGCACTTCCTCTGCTGTGTCTCCCACGAGGGGTGCCAGGTGCAGGCTCCAAAAGTA CTGGACGCAGAGGCCCTGCCCTTGCCAGACGGCGCCGAGACATGTCAGGACTAAGGAT CAGGCGCGCGTCGTCCTGCGTGTCTCTCCGAGGCGCCAGACATGTCAGGACAAAGTAAGGAT CAGGCGCGCGGTCCTGCTGTGTCTCTCGGAGGCGCCAGACATGTCAGGACAAAGTAAGGAT CAGGCGCGCGGTCCTGTCTGTGTCTCTGCGAGGCGCCAGACATGTCAGGACAAAGTAAGGAT CAGGCGCGCGGTCCTGTCTGTGTCTCAGACGGCGCCAGACATGTCAGGACAAAGTAAGGAC CAGGCGCGCGGTCCTGTCTGTGTCTCAAGACCCGAACATGTCAGGACAAAGTAAGGAC CAGGCGCCGCGGTCCTGTCTGTGTCCGAAGCGCCCCAACATGTCAGGACAAAGTAAGGAC CTGAGGGGCCCTAACTTTGCAGGCGCGCCCCAACATGTCGGAGGAGCACGAGGCCCTGGG CTGAGGGGCCCTAACTTTGCAGGCGCGCCCCAACTCCGAGGTGCACGAGGCCCCTGGG CTGAGGGGCCCTAACTTTGCGCGCGGACCCCGGACTGTGGGGGTGCACGGAGGCCCCGACGTGTGCGGGGTCCCCGGACGTGCGCGACGTGCGGGGGTCCCCGGACGTGCGGGGGTCCCCGACGTGCGGGGGGCCCCCAACTCCGGGGGCGCCCCAACTCCGGGGGGCCCCCGACGTGCGGGGGCCCCGACGTGCGGGGGGCCCCGGGGCGCCGGACGTGCGGGGGGCCCCGACGTGCGGGGGGCCCCGACGTGCGGGGGCGCCGGACGCGGGGGCCCCGGACGTGCGGGGGCCCCGGACGTGCGGGGGCGCCGGACGCGGGGGCCCCGGACGTGCGGGGGGCCCCGGCGACGCGGGGGCGCCGGCGCGGGCGCCGGGGGCCCCGGCGACGGCGG	3964 4566 $ 4566$ 5166 $ 5174$ 5166 $ 5174$ $ 5766$ $ 6294$ $ 6294$ $ -$	-39 -99 -99 -39 -39 -39 +21 +21 +21 +21 +21 +21 +21 +21 +21 +21
Mouse Rat JW HM HR MR · · · Human Mouse Rat JW HM MR · · · · · · · · · · · · · · · · · ·	ATGCGCAATATCGCATCAGCCCCCCCCG-CCCCGCTCCTCCCGGAAGGCCCAATC ATCCCCUARTATACCTTCGCCCCCCCCCCCCCCCCCCCCCCCC	3964 4554 $$ 4564 55164 $$ 516455 $$ 57745 $$ 5764355 $$ 62944 6695 $$ 62944 $$ 62944 $$ 62944 $$ $$ 62944 $$	-39 -99 -99 -39 -39 -39 -39 +21 +21 +21 +81 +81 +81 +129 +141
Mouse Rat. 3W HM HR HR MR Mouse Rat 3W HM HR HR Mouse Rat 3W HM HR MR HR HR MR HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTGCTGCGGAAGGCCCAATG ATGCGCAATATACGCTCTGCCCCGCCGCCGCCGTGCTGCGGAAGGCCCAATG CCCGCAGCGAACGCGCACTTCTCTCTCCGGAAGACTCCGCGGAGGGGGCGCGAGGGCCCGAAGAGCCC CTCAGTAAAGCGCACTTCCTCTGCGTAGTATCCACTGAGGGGGCGCAGGCCCGAGAGAGCC CTGGACGCAGAGGCCCTGCCCTGCCAGGCGCGCGAGACATGTCAGAAGAAAGTAAGGAT CAGGCGCGCAGTGGTCCTGCGTGCGCAGGCCCGAACATGTCAGAACAAAGTAAGGAT CAGGCGCGCAGGGCCCTGCCCTGCCAGGCGCGCAGACATGTCAGAACAAAGTAAGGAT CAGGCGCGCAGGGCCCTGCCCTGCCCAGGCGCCGAACATGTCAGAACAAAGTAAGGAT CAGGCGCGCAGGGCCCTGCCCTGCCCAGGCGCCGAACATGTCAGAACAAAGTAAGGAT CAGGCGGCCCTAACTTGCCAGAGGCCCCCAACTCCGAAGGTCAGGACAAAGTAAGGAC CTGAGCGACCCTAACTTGCAGCGGAGGCCCCCAACTCCGAGGTGCAGAGAAGTAAGGAC CTGAGCGACCCTAACTTTGCAGCGGAGGCCCCCAACTCCGAGGTGCAGAGAAGGAAG	3964 4556 	-39 -99 -99 -39 -39 +21 +21 +21 +21 +21 +21 +21 +21 +21 +21
Mouse Rat JW HM HR HR MR · · Rat JW HUMAN MR · · · HUMAN MOUSE Rat JW HUMAN HR HUMAN HR Kat JW HUMAN HR Kat JW HUMAN HR HIMAN HR · ·	ATGCGCAATATCGCATCAGCCCCCCCCG-CCCGCTCCTCCCGGAAGGCCCAATG ATCCCCUARTATACCTTC	3964 456 $ 456$ 514 516 $ 516$ 557 $ 5634$ $ 6695$ $ -$	-39 -99 -99 -39 -39 -39 -39 +21 +21 +21 +81 +81 +129 +141 +141
Mouse Rat. 3W HM HR HR MR HR Human Mouse Rat 3W HM HR Rat 3W HM HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCARTATCGCATCAGCCCCGCCCG-CCCGCTCCTCCGGAAGGCCCATG ATCCCGARTATACCGTC	3964 4556 $ 4566$ 5575 $ 5766355$ $ 5766355$ $ 6294$ 6695 $ -$	-39 -99 -99 -39 -39 -39 +21 +21 +21 +21 +21 +21 +21 +21 +21 +21
Mouse Rat 3W HM HR HR MR Human Mouse Rat 3W HM HR HR HR HR HR HR HR HR HR HR HR HR SW HM HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCAGCCCCCCCCGCCCCCCCCCC	3964 456 $ 456$ 516 $ 516$ 557 $ 576$ $ -$	-39 -39 -39 -39 -39 -39 -39 +21 +21 +21 +81 +81 +129 +141 +141
Mouse Rat JW Rat JW HM MR MR Mouse Rat JW HM HR MR Rat JW HM HR MR MR MR MR MR MR MR MR MR MR MR MR MR	ATGCGCAATATCGCATCAGCCCCGCCCGCCCGCCCCTCCCCGGAAGGCCCCATG ATGCGCAATATCGCATCAGCCCCGCCGCCGCCCGCTCCTCGGGAAGGCCCCATG ATGCGCAACACGCGCACTTCCTCCCGGGAATCCGCCGGAGGGGCGCGGGGGGGCCCGAAGGCCCCAATG CGCAGCAACGCGCACTTCCTCTCCGGGAATCCGCCGGAGGGGCGCGGGGGGGCCCCGAAGGCCC CTCAGTAAAGCGCACTTCCTCTCGGGAGTCCCCCCCGAGGGGCGCGGGGGGCCCCGAAGGCC CTCGGACGCACGCCCTGCCCTGCCCAGGCGCGCGGAGGACGCCCCGAAGGCTC CTGGACGCAGGGCCCTGCCCTGCCCAGCGCGCGCGCAGGCACGGCCCCGAAGGAC CTGGACGCGACGCCCTGCCCTGCCCAGCGCGCGCAGGCCCCGAAGGCCCCGGAGGACGCCCCGGAGGCC CTGGACGCGACGCCCTGCCCTGCCCCGACGCCCCGAACTCCGGAGGCACGGCCCCGGAGGCC CTGAGGCGACCCTAACTTTGCAGCGGAGGCCCCCGAACTCCGGAGGCACGGCCCCGGC CTGAGGCGACCCTAACTTTGCAGCGGAGGCCCCCGAACTCCGGAGGCACGGCCCCGGC CTGAGGCGACCCTAACTTTGCAGCGGAGGCCCCCGAACTCCGGAGGCACGGCACGCCTGGC CTGAGGCGACCCTAACTTTGCAGCCGGAGGCCCCCGAACTCCGGAGGCACGGCACGCCTGGC CTGAGGCGACCCTAACTTTGCAGCCGGAGGCCCCCGAACTCCGGAGGCACGGCACGCCTGGC CTGAGGCGACCCTAACTTTGCAGCCGGAGGCCCCCGAACTCCGGAGGCACGGCACGCCTGGC CTGAGGCGACCCTAACTTTGCAGCCGGAGGCCCCCGGACTGCGCGGCGCGCGC	3954 4556 $ 4516$ $ 5166$ 5775 $ 576435$ $ 576435$ $ 576435$ $ 57644$ $ 6294$ $ 6754$	-39 -99 -99 -39 -39 -39 -39 +21 +21 +21 +21 +81 +81 +129 +141 +141 +141 +189 +201
Mouse Rat 3W HM HR MR MR Mouse Rat 3W HM HR HR HUMAN HR HUMAN HR HUMAN HR HUMAN HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCAGCCCCCCCG-CCCGCTCCTGCGGAAGGCCCAATG ATGCGCAATATCGCATCAGCCCCCCCGCCGCCCCTCTCGCGAAGGCCCCAATG ATGCGCAACACGCGATCTGCCCCGCGGGGGCCCCGCCCC	3964 456 $$ 45635 $$ 5575 $$ 56335 $$ 66945 $$ 68844 7555	-39 -99 -99 -39 -39 -39 +21 +21 +21 +21 +21 +21 +21 +21 +21 +21
Mouse Rat 3W HM HR HR MR · · Human Mouse Rat 3W HM HR Max · · · Human Mouse Rat 3W HM HR · · · · · · · · · · · · · · · · · ·	ATGCGCAATATCGCATCAGCCCCCCCCG-CCCCGCCCCCCCCGCGAGGCCCCATC ATCCCCUARTATCACCTC	3964 456 $ 4514$ 516 $ 5774$ 575 $ 5764635$ $ 6294$ 6754 $ 62945$ $ 6844$ $ -$	-39 -99 -99 -39 -39 -39 -39 -39 +21 +21 +21 +81 +81 +141 +141 +141 +189 +201 +201
Mouse Rat 3W HM HR MR MR Mouse Rat 3W HM MR HR MR HR MR HR MR HR MR HR MR HR MR HR MR HR MR HR HR MR HR HR MR HR HR MOUSE Rat 3W HM HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCAGCCCCGCCCG-CCCGCTCCTCCGGAAGCCCCAATG ATCCCCAATATACCGTTGCCCCGCCCGCCCGTCTCCGGAAGCCCCAATG CCCCCCAATATACCGTTCTTCTCTCCGGCAGAATCCGCGGAGGGCCCGGAGGGCCCGAAGGCTC CTCAGTAAAGCGCACTTCTTCTCTCCGGCAGCGCCGGAGGGCCGCGGGCCCGAAGGCTC CTCAGTAAAGCGCACTTCCTCTGCTAGTATCCACTGAGGGCCCGGAGGGCCCGAAGGCTC CTGGACGCAGAGGCCCTGCCCTGCCCAGGCGCGCGGACATGTCAGAGGCTCCGAGAGGTC CTGGACGCAGAGGCCCTGCCCTGCCCAGACGGCGCGGACATGTCAGAGCAAAGTAAGGAT CAGCGGCGCCTGCCTGTCGTCGCCGGAGGCCCCGAACATGTCAGAACAAAGTAAGGAT CAGCGGCGCCTGCCTGCCCTGCCCAGGCCGCGAGACATGTCAGAACAAAGTAAGGAC CTGGACGCAGAGGCCCTGCCCTGCCTGCGCAGGCCCCGAACATGTCAGAACAAAGTAAGGAC CTGGACGGACCGTAACTTTGCAGCGGAGGCCCCCAACTCCGGAGGACGAAGGAAG	3964 4456 $ 4514$ 5516 $ 5164$ 5775 $ 56635$ $ 62945$ $ 6884$ 854 $ 6854$ $ -$	-39 -99 -99 -39 -39 -39 +21 +21 +81 +61 +61 +61 +81 +129 +141 +141 +141
Mouse Rat JW HM HR HR MR · · Human Mouse Rat JW HM MR · · Human Mouse Rat JW HUMAN HR HR HR HR HR HR HR HR HR HR HR HR HR	ATGCGCAATATCGCATCAGCCCCCCCCG-CCCCGCTCCTCCGGAAGGCCCAATG ATCCCCUARTATACCTCTGCCCCCGCCGCCCCCTCCTCGGAAGGCCCAATG 	3964 456 $ 4516$ $ 557$ $ 5775$ $ 576635$ $ 6695$ $ 684$ 7555 $ -$	-39 -99 -99 -39 -39 -39 -39 -39 +21 +21 +21 +81 +81 +129 +141 +141 +141 +141 +141

Figure 3-1. Sequence alignment of *NDN/Ndn*.

ClustalW alignment of human, mouse and rat NDN/Ndn. Multiple alignment with all three sequences is shown. Asterisks indicate sequence identity across all three (3W), human versus mouse (HM), human versus rat (HR), and mouse versus rat (MR). Red asterisks indicate approximate location of identity in another frame with respect to the multiple alignments. The start codon is indicated in blue, the transcription start indicated in green, the minimal promoter between single and double daggers, and the two opposing arrowheads indicates division between weak and strong sequence conservation between human and mouse.

Figure 3-2. Percentage identity plot and predicted TF sites.

(A) Percentage identity plots for the upstream region of *NDN/Ndn* in human, mouse and rat. The Y-axis indicates percentage identity across 50 bp windows. X-axis indicates the position, numbered with start codon as +1. (B) Schematic showing features upstream of *NDN/Ndn*. Dark green rounded bar indicates minimal promoter. Green arrow indicates transcription start. Light green rectangular bar indicates ORF. Vertical black line indicates position -133, a division between weak and strong sequence conservation between human and mouse. Foot indicates the transcription-related *in vivo* footprint in human *NDN*. Small coloured circles indicate position of consensus binding sites for transcription factors found across all three species and at the same position.



The sequence of the transcription-related footprint characterized by Dr. Hanel was analyzed with MatInspector to find putative transcription factor binding consensus sites (Quandt et al. 1995). Within this region are consensus sites for MAZR (MYC-associated zinc finger protein related transcription factor, ZNF278) (Kobayashi et al. 2000) and a pair each of SP1 and NRF1 (Scarpulla 2002) sites that are conserved with mouse and rat with respect to position. Specific to the human upstream sequence are also additional NRF1, AP2, PAX5, SP2, EGR1, and CPBP sites (Figure 3-3) (Koritschoner et al. 1997; Busslinger 2004; Simon et al. 2004).

To further characterize conservation of sequence motifs that may be indicative of transcription factor binding sites, sequences upstream of *NDN/Ndn* were analyzed. Numerous sequences with similarity to transcription factor consensus sites were found in the human, mouse and rat sequences (112, 103, and 88 respectively). Putative factor binding sites significant to regulation of *NDN/Ndn* are expected to be conserved across all three species (Table 3-1). Furthermore, conservation in position would also be suggestive of conservation of an important factor binding site (Figure 3-2 B). Consensus sites for nine transcription factors were found in the promoter sequences of all three species, and in a similar position with respect to the start codon. Of the nine putative factors, there were two closely spaced consensus sites each for NRF1 and SP1 (Figure 3-3).

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Match summ	ary		
Family	Human	Mouse	Rat
BCL6	1	1	2
DEA-F	2	2	1
EGR-F	8	3	3
EKL-F	2	1	1
NRF2	1	1	1
ETS1	1	2	1
FKHD	1	2	1
HOM-F	2	2	2
HOX-F	1	4	2
LEF-F	1	2	1
MAZ-F	2	1	1
NO765	2	3	3
MYT1	2	4	4
NFKB	4	3	1
NKXH	3	2	1
NRF1	6	2	2
OCT1	1	1	2
RBP-F	1	1	2
RORA	2	1	2
SP1-F	6	2	1
TBP-F	1	2	1

Table 3-1. Table of predicted TFs.

Summary of putative transcription factor sites shared between human, mouse and rat. Numbers indicate multiples of those sites found in a particular species. Colored background corresponds to factors in Figure 3-2 B. Transcription-related footprint



NDN PAX5 oligo CCSCGGTCGCGCAGGC GCAGTGCCGCGTCCCG

NDN CPBP oligo Teccecce ceccececetece tecce

Figure 3-3. Predicted factor binding sites in transcription-related footprint.

Shown at top is the DNA duplex of positions -196 to -115 upstream of human *NDN* start codon. In blue, is the transcription-related footprint found on the expressing allele in the expressing tissue. Below are the same sequences, with putative transcription factor binding sites shown in red.

Verification of bioinformatically predicted PAX5 site by EMSAs

While binding sequences for NRF1 are coincident with the transcription-related footprint, making it a possible candidate for a factor occupying the footprint, the presence of these sites was not known until the latest update of the MatInspector database (version 7.4, Jan, 2005). The experiments described here are based on data from a previous database revision (6.1, Jan, 2003). The next best candidate for a factor responsible for the footprint is PAX5. Other than the two consensus sites found near the footprint, the human sequence has a third consensus site at position -216. To test if PAX5 can indeed bind the predicted sequence upstream of NDN, EMSAs were performed with the aid of Gareth Cory of the laboratory of Dr. Alan Underhill. Radiolabeled ds-oligonucleotides were incubated with proteins of interest to form protein-DNA complexes, which were detected after size separation on polyacrylamide gel electrophoresis. PAX5 binds DNA through its paireddomain, and has highly similar binding specificity to the other members of its subfamily, PAX2 and PAX8; therefore paired-domains from these proteins were used to test binding to upstream regions of NDN (Figure 3-4). These domains were produced in bacteria from constructs that also encoded a 6XHIS tag for purification purposes. The binding of these purified Pax2 and Pax8 paired-domains to double-stranded DNA representing the promoter region of NDN found to be occupied by the transcription-related footprint (positions -170 to -139, NDNPAX5) was tested. Both the Pax2 and Pax8 paired-domains were able to bind this 30 bp double stranded oligonucleotide NDNPAX5, although not as well as to an idealized consensus sequence of the same length bound by the PAX2/5/8 subfamily of paired-domains (PAX258, (Czerny and Busslinger 1995)) (Figure 3-4). The Pax2 paired

domain also had a qualitatively higher affinity to the NDNPAX5 sequence than Pax8 paired domain.



Oligo: PAX258 NDNPAX5 PAX258 NDNPAX5 Protein: Pax2 PD Pax8 PD

Figure 3-4. EMSA of footprinted site to related paired domains.

EMSA of labeled oligos made to transcription-related footprinted region in *NDN* promoter binding to Pax2 and Pax8 paired domains, which are highly related to the Pax5 paired domain. Oligo PAX258 is of an idealized binding sequence, and NDNPAX5 is of a 30b sequence occupied by the transcription-related footprint.



Figure 3-5. RT-PCR of *PAX* **genes in fibroblast lines.** RT-PCR of *PAX2*, *PAX5* and *PAX8* in PWS and AS patient fibroblasts (FB16 and FB17, respectively).

In humans, these proteins are normally expressed postnatally only in retina and kidney (*PAX2*) (Eccles et al. 2002; Pichaud and Desplan 2002), B-cell lineages (*PAX5*) (Borson et al. 2002) and thyroid (*PAX8*) (Christophe 2004). To further investigate the possibility that one of the *PAX2/5/8* subfamily of transcription factors may be expressed in cells where the footprint was found, we performed reverse transcription-PCR for these three PAX genes in the fibroblast cell lines used in the *in vivo* footprinting and found no evidence for expression of *PAX2, PAX5* or *PAX* 8 (Figure 3-5). This transcription-related footprint may thus identify a binding site for another human transcription factor such as NRF1, or an unidentified one related in binding specificity to proteins of the PAX family.

Allelic differences in footprints are due to chromatin states

It was necessary to determine if allelic footprints were due to differences between alleles in terms of chromatin accessibility or due to differences in complement of proteins capable of binding in the individual cell lines. Therefore, I tested if similar proteins are in each cell line that has the *in vitro* potential to bind the promoter sequences. Nuclear extracts were prepared from PWS and AS deletion fibroblasts and lymphoblasts and binding to *NDN* upstream sequences was assayed. The double-stranded oligonucleotide NDNPAX5 corresponds to the transcription-related footprint found on the paternal expressed allele of the fibroblast, while the ds-oligonucleotide NDNCPBP corresponds to a region containing the predicted CPBP, EGR-1 and MAZR sites (position -148 to -110) (Figure 3-2). Dsoligonucleotide NDNMOK corresponds to a region (+5 to +40) coinciding with a maternal allele-specific DNase I footprint found in both fibroblasts and lymphoblasts, and is coincident with a predicted MOK-2 consensus site. Similar to previous EMSA experiments, radiolabeled ds-oligonucleotides were incubated with nuclear extracts and size separated to give patterns indicative of protein-DNA complexes. As expected, we found no evidence that there are differences in the complement of factors able to bind the oligonucleotides between PWS and AS cell lines (Figure 3-6). This indicates there are factors available in the PWS fibroblasts that can bind to the footprinted sequence, but do not, presumably due to inaccessibility of the chromatin on the maternal allele. DNA methylation may also play a role here to exclude binding of factors, as it has been shown that the *NDN* promoter is more methylated on the maternal allele in this region (Lau et al. 2004).

Figure 3-6. EMSAs with patient cell line nuclear extracts.

EMSAs with nuclear extracts from PWS and AS fibroblasts and lymphoblasts. Pax2 paireddomain (PAX2pd) included for comparison. NDNPAX5 corresponds to the transcription-related footprint in AS fibroblasts, NDNCPBP corresponds to predicted factor sites downstream of the footprint, and NDNMOK corresponds to MOK-2 putative site downstream of the promoter. For each pair of EMSA reactions, the left and right lanes indicate addition of 5µl and 10µl of extract, respectively.







To investigate the presence of specific candidate factors that may bind the *NDN* promoter in both our fibroblast lines, expression of *CPBP*, *EGR-1*, *MAZR* and *MOK-2* were tested by RT-PCR (Figure 3-7). We found no evidence that there are differences in the expression of candidate factors between PWS and AS cell lines. This is consistent with the nuclear extract EMSA data above, which does not show any differences in binding patterns of oligonucleotides between PWS and AS cells. In contrast to the *PAX2/5/8* family, *CPBP* and *EGR-1* are strongly expressed in both fibroblast lines. *MOK-2* is expressed at limited levels, while *MAZR* expression is not detectable.



Figure 3-7. RT-PCR of candidate factors binding to the *NDN* promoter.

RT-PCR of candidate binding factors in PWS and AS patient fibroblasts.

Discussion

The regulation of *NDN* is influenced by multiple factors including chromatin state and *trans*-acting factors that direct its correct allelic and developmental expression. The *in silico* approach to predict possible transcription factor binding sites and the complementary *in vivo* data from the work of Dr. Meredith Hanel has given insight into a cross-section of these layers. The bioinformatics analysis has shown several putative conserved factor binding sites in regions of high cross-species sequence conservation, and surprisingly, also in regions of little conservation between human and rodent (Figure 3-2). Furthermore, investigation of the *in vivo* occupancy of the human *NDN* promoter by footprinting has aided identification of potentially important regulatory elements without cross-species conservation. Similar approaches have led to functional characterization of factors regulating other genes (Smith et al. 2004b).

The bioinformatic prediction of transcription factor binding sites suggested several motifs were found across human, mouse and rat, and in a similar position relative to the open reading frame of *NDN/Ndn*. Each transcription factor may have a role in regulating the tissue or allele-specificity of transcription. MyT1 is a transcription factor that is thought to interact with neurogenin 1 and specifies vertebrate early neuronal precursor fates by affecting the *Notch* signaling pathway (Quan et al. 2004). This is consistent with a role for *NDN* in brain function (Ren et al. 2003; Lee et al. 2005). It is also notable that there are multiple possible MyT1 sites found in human, mouse and rat, one of which is conserved in position to *NDN/Ndn* in a region of little overall sequence conservation, which suggests it may be a functional component of the regulatory elements controlling *NDN/Ndn*.
NRF-1 and SP1 sites are also found upstream of *NDN/Ndn* in all three species. NRF (nuclear respiratory factor) proteins function to regulate nuclear genes important in mitochondrial biogenesis and function (Scarpulla 2002). While *NDN* has no previously characterized role in mitochondrial function or energy metabolism, NRF-1 has been shown to be involved in regulation of *FMR1*, a gene that has also not been shown to be immediately involved in mitochondrial function (Smith et al. 2004a). In fact, at the *FMR1* promoter, NRF-1 synergistically interacts with SP1, whose binding sites are also predicted to be in close proximity in the promoter of *NDN*. It will be of interest to investigate whether the same interaction and activation between NRF-1 and SP1 also occurs to regulate *NDN*.

Little is known about the function of MAZR, although it interacts with Bach2, a Bcell and neuron-specific transcriptional repressor (Kobayashi et al. 2000). A neuronspecific function would be of interest to *NDN* biology, but a role in B-cells has yet been suggested for *NDN*. Like MAZR, putative sites for LEF1 are also found in all three species. LEF1 is a factor involved in bone morphogenesis in the Wnt signaling pathway (Westendorf et al. 2004), as well as response to survival signals in pro-B-cells (Busslinger 2004). There are also conserved putative binding sites for ETS-1, which is a protooncogene that has been shown to interact with many other transcription factors, one of them being PAX5, to regulate B-cell differentiation and immunoglobulin heavy chain expression (Dittmer 2003). It was therefore of special interest that two PAX5 binding sites were predicted overlapping the transcription-related footprint found on the expressed allele in fibroblast, possibly indicating a complex important for the transcription of *NDN*. PAX5 plays a central role in specifying B-cell identity (Busslinger 2004) and has been shown to be involved in the histone exchange removal of H3K9 methylation during VDJ rearrangement (Johnson et al. 2004). Our data shows that while the highly related Pax2 and Pax8 paired-domains are able to bind *NDN* upstream sequences *in vitro* (Figure 3-4), PAX5 itself is not detectably expressed in our fibroblast lines (Figure 3-5), nor are the sites conserved with rodents. *NDN* is also not expressed in blood lymphocytes, which would include cells of the B-cell lineage. Therefore a link between *NDN* and B-cell function is not supported, but cannot be ruled out in other cell types.

CPBP and *EGR-1* are expressed in the fibroblast cells in which the footprinting assay was performed, and as such, remain candidates for factors binding at the footprint. While the position of their predicted binding sites are not completely overlapping the *in vivo* footprint, their binding and associated complex may change local DNA structure and change sensitivity to DNase I, DMS and UVC modification. CPBP (core promoter binding protein) is a ubiquitous factor that functions in many genes, more specifically, at TATAless promoters (Koritschoner et al. 1997) like the promoter of *NDN*. EGR-1 is an immediate-early gene that is involved in responses to a variety of signaling cascades that modify transcription of target genes. More specifically EGR-1 has been implicated in synaptic plasticity (Simon et al. 2004) and oncogenesis (Adamson et al. 2005). Whether EGR-1 participates in the same pathways as *NDN* does in brain development remains to be investigated.

Putative binding sites for MOK-2 are conserved between all three species. With respect to the human footprinting data, one of the MOK-2 sites overlaps a DNase I hyposensitivity region on the maternal allele in both fibroblasts and lymphoblasts, making

Introduction

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As discussed in the Introduction, epigenetic processes such as DNA methylation and histone modifications are associated with many processes such as transcriptional regulation (El-Osta and Wolffe 2000; Litt et al. 2001; Noma et al. 2001), X-inactivation (Beard et al. 1995; Keohane et al. 1998; Boggs et al. 2002) and genomic imprinting (Brannan and Bartolomei 1999; Mann et al. 2000; Grandjean et al. 2001; Xin et al. 2001). Mechanistic links between DNA methylation and histone modification have been proposed, whereby histone H3 K9 methylation can direct DNA methylation (Rice and Allis 2001; Tamaru and Selker 2001), which can then in turn recruit histone deacetylases, thereby creating a closed chromatin conformation that inhibits transcription (Nan et al. 1998; El-Osta and Wolffe 2000; Jones and Takai 2001). DNA hypermethylation is generally associated with decreased gene activity (Jones and Takai 2001; Reik et al. 2001). Histone hyperacetylation is associated with actively transcribed genes and genes poised for transcription (Razin 1998). Histone H3 can be methylated on lysine 4 and lysine 9 residues

strikingly different (Figure 3-6), suggesting tissue-specific lack of footprints may also be in part due to chromatin differences instead of a lack of *trans*-acting factors between fibroblasts and lymphoblasts. Further investigation of the differences in chromatin structure and composition of the two alleles, as presented in the following chapters, will clarify the state of the chromatin at *NDN/Ndn* and how this regulates its allele-specific and tissue-specific transcription.

Chapter 4 ♦ Tissue-specific and imprinted histone modifications of the human NDN gene

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Introduction

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to mark either active or inactive chromatin (Jenuwein and Allis 2001; Noma et al. 2001). Whether the same mechanisms operate in tissue-specific control and allele-specific control is less well understood.

The *SNRPN* gene, located in the Prader-Willi Syndrome (PWS) region on 15q11q13, and the *H19/IGF2* gene pair are among the most intensively studied imprinted genes (Ferguson-Smith and Surani 2001). These genes contain imprinting control elements that control germline imprint resetting of genes located in *cis*, even over large distances (Leighton et al. 1995; Dittrich et al. 1996; Horsthemke 1997). Allelic epigenetic differences found at these imprinted loci with closely associated Imprinting Centers (IC) can therefore be either associated with the IC itself, or be the result of a response to the IC (Pedone et al. 1999; Schweizer et al. 1999; Saitoh and Wada 2000). Current imprinting models do not address mechanisms for the extension of the epigenetic mark to target genes at a distance from their IC, nor the mechanisms for coordinate allele- and tissue-specific expression (Hu et al. 1998; Hanel and Wevrick 2001).

As described in the rationale in Chapter 3, *NDN* serves as an excellent model for the transposed imprinted genes. Our laboratory has previously demonstrated developmentally dynamic patterns of maternal hypermethylation and paternal hypomethylation of the promoter CpG island in mouse *Ndn*, by sodium bisulfite sequencing (Hanel and Wevrick 2001). In this chapter, I will describe characterization of finely mapped regions of histone acetylation and histone methylation surrounding *NDN* using antibody specificities previously shown to be differentially modified in imprinted regions. In contrast to *SNRPN*, *NDN* has a tissue-specific expression pattern and is

expressed in brain and fibroblasts among other tissues but is silent in blood lymphocytes and derivative lymphoblastoid cell lines (Jay et al. 1997; MacDonald and Wevrick 1997; Sutcliffe et al. 1997; Hanel and Wevrick 2001). This allows comparisons of DNA methylation and histone modification between tissues in which *NDN* is and is not expressed. The simple intronless genomic structure of *NDN* is also amenable to high resolution ChIP to study the histone modifications over the entire transcription unit. These data were compared with complementary experiments by Dr. Meredith Hanel on the allelespecific methylation of a promoter CpG island and a second downstream CpG island in human *NDN* by sodium bisulfite sequencing. We have evaluated the relative contribution of epigenetic changes associated with tissue-specific gene expression versus those associated with genomic imprinting. Our results suggest that DNA methylation and histone H3 K4 dimethylation and trimethylation epigenetically differentiate alleles in *NDN*, while histone acetylation acts in tissue-specific gene regulation.

Results

Histone modification of NDN.

Chromatin immunoprecipitation (ChIP) was performed to investigate histone modifications across the *NDN* gene. In order to analyze the maternal and paternal alleles in isolation, we again used cell lines derived from AS individuals carrying maternal deletion of 15q11-q13, or PWS individuals carrying a paternal deletion. Primer sets were designed to give high resolution coverage (Figure 4-1). Regions B-G covers the transcription unit of

NDN and 5' CpG island, while regions A and H are several kilobases upstream and downstream respectively. We first analyzed fibroblasts derived from PWS and AS patients. Consistent with previously identified patterns of histone H3 acetylation (H3ac), we observed paternal bias in NDN in all regions assayed inside and outside of the transcription unit (Figure 4-3). A similar paternal bias in H4 acetylation was also present (Figure 4-2). While differences in acetylation were present across NDN, consistent allelic differences were largest in region B, colocalizing with the promoter where there were greater than four-fold differences between alleles. We then performed similar ChIP analysis with antibodies specific for di- and tri-methylated forms of lysine 4 and lysine 9 of the histone H3 tail (H3K4me2, H3K4me3, H3K9me2, and H3K9me3, of which, H3K9me3 cross reacts with H3K27me3 (Perez-Burgos et al. 2004)) and di-methylated lysine 79 of the histone H3 globular domain (H3K79me2). Consistent paternal bias in H3K4me2 was observed over regions B-E (Figure 4-3). The most striking H3K4me2 difference was seen in region B with an average of greater than seven-fold paternal bias. Using trimethyl specific antibodies, a more restricted pattern of paternal bias in H3K4me3 was seen consistently over region B only. H3K4me3 showed approximately three-fold paternal enrichment with very weak or inconsistent biases elsewhere in the gene. Results characterizing H3K9me2 were quite inconsistent, with qualitative maternal biases in some early experiments (Figure 4-2), but a paternal bias in certain regions in others (Figure 4-4). The H3K79me2 and H3K9me3 (and H3K27me3 by cross reactivity of the antibody) antibodies detected a variable and weak trend towards paternal bias (Figure 4-4).

Unambiguous analysis of this modification awaits commercial availability of more specific antibodies. More detailed investigation of these modifications was not pursued further.





Figure 4-1. NDN and surrounding regions.

Dark green box indicates the single exon of the *NDN* gene, with the ORF indicated by the inset light green box. The arrow indicates transcription start site and CpG islands are as indicated. Regions analyzed with in vivo footprinting in Chapter 3, and bisulfite sequencing by Dr. Meredith Hanel are indicated by "LM-PCR", and double-headed arrows, respectively. Regions analyzed by ChIP are indicated by amplicons A through H. The two gaps each represent approximately 3 kb of DNA. Orange region indicates extent of human-mouse homology.



Figure 4-2. Examples of qualitative and quantitative ChIP analysis.

(A) Qualitative analysis of H3ac, H4ac, H3K4me2, and H3K9me2 in PWS (left band in each frame) and AS (right band in each frame) fibroblast and lymphocytes. Note the apparent maternal bias in H3K9me2 in amplicon E. (B) Quantitative analysis of band intensities by phosphorimager scanning of radiolabeled probes annealed to slot-blotted PCR products. N* indicates a region overlapping with N. Lower right inset is an example of the quantitation object drawn around a band in the ImageQuant software.



Figure 4-3. Histone modifications of active transcription.

Quantitative ChIP data from experiments using antibodies against H3ac, H3me2K4, and H3me3K4 in PWS and AS fibroblast and lymphocyte data is shown. Paternal to maternal ratio of a representative trial plotted on logarithmic scale where one indicates no bias, greater than one is a paternal bias and less than one is maternal bias. Letters correspond to amplicons assayed as described in (Figure 4-1). Shown is a representative trial, with error bars indicate variation of multiple rounds of detection.



Figure 4-4. Preliminary quantitative data on ChIP in PWS and AS fibroblast lines.

Antibodies against H3K79me2, H3K9me2 and H3K9me3 were used for the experiments below. Note that the paternal bias in H3K9me2 is not always observed (Figure 4-2).



Figure 4-5. ChIP data with antibodies against various histone modifications at SNRPN. Data presented as in Figure 4-3.

We next performed similar experiments in patient blood lymphocytes to assay whether or not the H3K4me2 and H3K4me3 paternal biases were correlated with tissue type and NDN expression. It was previously reported that a region within region F is not associated with allelic histone acetylation in lymphoblasts (Fulmer-Smentek and Francke 2001) and the paternal allele is associated with histone H3K4me2 in region C in blood lymphocytes (Xin et al. 2001). We confirmed this lack of allelic histone acetylation in PWS and AS LCLs and lymphocytes in region F as well as paternal H3K4me2 of region C in patient blood lymphocytes. No other regions in NDN had consistent allelic histone acetylation in lymphocytes (Figure 4-2). H3K4me2 allelic differences were distributed over a wider region than previously reported, covering most regions analyzed, although with a weaker bias than seen in fibroblasts (Figure 4-3). A trend towards paternal enrichment for H3K4me3 was also found in lymphocytes, although the degree and distribution of this bias was much more variable. Other modifications were not investigated in lymphocytes because only a limited amount of patient material was available. Overall, these results define a domain of paternal H3K4me3 lying within a domain of paternal H3K4me2 which itself is contained within a large domain of paternal H3ac in fibroblasts, while lymphocytes show a more general allelic bias in H3K4me2 and H3K4me3 without allelic H3ac.

Histone modification of the IC.

To make comparisons between *NDN* and its imprinting center, we studied *SNRPN/IC*, which is expressed in fibroblasts and lymphocytes. We examined histone modification in exon 1 of *SNRPN*, previously described to be paternally enriched for histone H3K4me2 and maternally enriched for histone H3K9me2 in lymphocytes (Xin et

al. 2001). In fibroblasts, an H3me2K4 paternal bias was also seen, while we observed maternal bias in H3me2K9 in only some of our trials (Figure 4-5). We next determined if this bias extended to the trimethylated forms, H3K4me3 and H3K9me3. H3K4me3 was found to be paternally enriched at *SNRPN* exon 1 at a level comparable to the enrichment seen with H3K4me2 (Figure 4-3). Using antibodies specific to H3K9me3 however, did not show significant differences between alleles. In blood lymphocytes, we confirmed the paternal bias previously seen in H3K4me2, and discovered an H3K4me3 bias, as is seen in fibroblasts (Figure 4-5). Similar to *NDN*, only weak and inconsistent biases were seen in H3K79me2. These observations are consistent with the fact that *SNRPN/IC* is expressed from the paternal allele in both fibroblasts and lymphocytes.

Discussion

Concurrent with these studies on histone modifications at *NDN* and *SNRPN*, Dr. Meredith Hanel performed bisulfite sequencing to determine the methylation status of *NDN*. She identified a 5' CpG island of 880 bp, containing 73 CpG sites, located in the promoter region of *NDN*, extending from 335 bp upstream of the start codon, and into the open reading frame (Figure 4-1). A second CpG island is located about 4.3 kb downstream of the *NDN* start codon; no equivalent downstream CpG island was found in the mouse sequence for 30 kb downstream of *Ndn* (Genbank #AC027298). Overall, DNA hypermethylation in the 5' CpG island of the maternal allele compared to the paternal allele was observed in both fibroblasts and blood lymphocytes (Lau et al. 2004). The 3' CpG island carried no allelic DNA methylation patterns. Our studies of histone acetylation are consistent with findings that developmentally regulated genes, such as *NDN*, are usually associated with domains of hyperacetylation (Hebbes et al. 1994; Forsberg et al. 2000) while changes in gene activity in response to stimuli are more frequently associated with localized changes in acetylation (Parekh and Maniatis 1999). In fibroblasts, in which *NDN* is actively transcribed, we identified allelic acetylation differences in a region of at least 10 kb surrounding *NDN*. Intriguingly, the paternal allele is hypoacetylated in the absence of DNA methylation in lymphocytes, suggesting that at this locus histone deacetylases are recruited by factors that are not dependent on DNA methylation, or that DNA methylation is lost after establishment of the hypoacetylated state. Limited studies of the murine *Ndn* promoter, in a region equivalent to human region D (Figure 4-1), indicate that neither allele is acetylated in liver, where necdin is inactive, whereas at least one allele is acetylated in brain, where necdin is expressed (Forsberg et al. 2000). Thus in both human and mouse, the acetylation state of *NDN* may act transiently in transcriptionally competent and transcriptionally active cells, and does not appear to remain as a longer lasting epigenetic imprinting mark.

In *S. cerevisiae*, H3K4me2 has been associated with euchromatic regions of the genome whereas a H3K4me3 state is only seen in actively transcribed genes (Litt et al. 2001; Noma et al. 2001). In light of this association for H3K4me3, it is of great surprise that lymphocytes, not actively transcribing *NDN*, would carry any paternal bias in this modification. We observed a paternal bias in H3K4me2 and H3K4me3 at *NDN* in lymphocytes (Figure 4-3). This shows a striking resemblance to the β -globin cluster in that inactive β -globin genes still carry H3K4me2 and H3K4me3 modifications, which is in

contrast to other developmentally regulated genes (Schneider et al. 2004). As those authors suggest, one possible explanation may be related to the long range function of the β -globin LCR, and at this locus, the PWS/AS IC may share similar mechanisms of action. It is possible that the maintenance of the paternal state within the PWS/AS cluster requires all genes on that allele carry certain epigenetic marks regardless of tissue-specific transcriptional status.

Of greater interest are the wide region of paternal H3K4me2 and the nested region of H3K4me3 in fibroblasts. These modifications have been found to be markers of euchromatic regions and transcribed genes respectively (Litt et al. 2001; Noma et al. 2001). Unlike histone acetylation, histone methylation status has been implicated as an early event in chromatin control (Rice and Allis 2001), with histone H3 methylation on residues lysine 4 and lysine 9 reciprocally marking active chromatin and heterochromatin respectively (Jenuwein and Allis 2001). As allelic H3K4me2 and H3K4me3 is also present in lymphocytes (Xin et al. 2001), we propose a model whereby histone H3 methylation at lysine 4 acts to mark allelic differential chromatin states at the NDN locus in response to the IC, and that this histone modification represents a persistent somatic mark of the active allele that allows histone acetylation to regulate expression of NDN in a tissue-specific manner (Figure 4-6). Interestingly, it has recently been shown that a promoter-restricted distribution of H3K4me2 is a marker of monoallelic genes (Rougeulle et al. 2003). While these authors were not able to distinguish parental alleles, we show here that a similar bias is present in NDN on the paternal allele regardless of expression. Restriction of the H3K4me2 modification to near the promoter of this single exon gene is also consistent with

their observations. It remains to be seen whether H3K4me3 also display distribution patterns characteristic of imprinted or other monoallelic genes versus biallelic genes. The multiple levels of histone modification in expressing and non-expressing tissues and persistent allelic identity of this imprinted locus may indicate involvement of remodeling complexes implicated in cellular memory. For example, the human trithorax group ALL-1 complex contains HMT activity towards H3K4, HAT activity, as well as chromatin remodeling activity (Nakamura et al. 2002). It will be of great interest to study association of this or other regulatory complexes to maintenance of imprinting at the PWS/AS cluster.



Figure 4-6. Model in which histone methylation plays an early role in imprint establishment and maintenance in NDN.

Pat and Mat refer to alleles of paternal and maternal origin. Black arrow indicates transcription of *NDN* on the paternal alelle in expressing tissue. Lollipops on histone (cylinders) residues and spiral DNA strand indicate allelic biases in histone lysine methylation and DNA CpG methylation respectively. Triangles indicate histone H3 acetylation differences. Solid black symbols indicate bias in allelic modification of that epigenetic mark. The initial signal from the imprinting center can determine histone methylation states, which is translated into other epigenetic marks such as DNA methylation and histone acetylation which sets up a chromatin context for DNA binding factors.

The developmental origins of somatic maternal epigenetic marks are not clear. In mouse and human oocytes, the *NDN/Ndn* promoter is variably methylated, and at least in mouse, differences between the parental alleles are no longer present in blastocysts (El-Maarri et al. 2001; Hanel and Wevrick 2001). Histone H3 K4 methylation on the active allele could serve as a candidate initial epigenetic mark of imprinted target genes, or could translate an initial DNA methylation imprint into a long-term mark that differentiates the two alleles. It remains to be tested whether differential histone H3 methylation exists during early embryogenesis and throughout development and if it acts at the top of a chromatin control hierarchy above allelic DNA methylation and histone acetylation or is simply correlated with these other epigenetic differences. While the mechanism of chromatin changes at target genes by the imprinting center is unknown, our data suggests that allele-specific and tissue-specific epigenetic changes are coordinated for proper gene expression.

Chapter 5 CTCF binds differentially methylated regions in the imprinted mouse Prader-Willi Syndrome locus

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Introduction

Chapter 4 described the histone modification profile in the transcription unit of *NDN*, a representative transposed imprinted gene (Figure 4-3). It is important to study how these modifications apply to the other genes, as well as the conservation of these modifications in other organisms to infer biological significance. Also, since the transposed imprinted genes are co-regulated, it is of interest to investigate mechanisms of domain-wide allelic and tissue-specific regulation of this cluster. The mouse system offers an ideal model to study both these aspects. Mouse chromosome 7C harbors an imprinted domain that is 3.3 Mb in size, and has conserved synteny with human chromosome 15q11-13, the PWS/AS region (Nicholls and Knepper 2001). As described in the Introduction, parental gender of origin-specific gene expression is under the control of the PWS/AS IC located upstream of *SNURF-SNRPN*, which carries a germline maternally DNA methylated region (DMR), paternal histone H3 lysine 4 (H3K4me2) methylation, and maternal H3 lysine 9 (H3K9me2) methylation (Xin et al. 2001). Histone methylation of K4 and K9 associated with the regulation of gene activity with methylation of K4 and K9 associated with activity and silence respectively (Peterson and Laniel 2004).

The imprinting of genes located centromeric to the IC on mouse chromosome 7C is mediated by a Megabase long RNA transcript produced from the paternal allele, which controls the paternal allele-specific expression of the contiguous genes *Snurf-Snrpn*, *snoRNAs*, and also the maternal allele-specific expression of *Ube3a* and *Atp10c* through an antisense mechanism (Chamberlain and Brannan 2001; Runte et al. 2001; Landers et al. 2004). However, there is no indication that a similar RNA-based mechanism maintains

imprinting for the contiguous genes *Frat3*, *Mkrn3*, *Magel2* and *Ndn*, which are located together on the telomeric side of the IC and all transcribed from the paternal allele. Unlike the *Snrpn-Atp10c* cluster, which are transcribed telomere to centromere for paternal genes and reverse for maternal genes, the paternally expressed *Frat3-Ndn* genes are transcribed centromere to telomere for *Ndn* and *Magel2*, and reverse for *Mkrn3* and *Frat3* (Figure 5-1, top), making an antisense RNA mechanism less plausible. These genes, referred to as the transposed imprinted genes, are thought to have been evolutionarily recent additions to the gene cluster and are proposed to have become imprinted as bystanders (Chai et al. 2001). There is a no homolog for *Frat3* in the human PWS/AS region, where instead the *Mkrn3* homologue *MKRN3* is the most distally located imprinted gene in the PWS/AS cluster.

We therefore investigated other epigenetic mechanisms that could be responsible for maintaining the imprinting status of the transposed imprinted genes. At the *H19/1gf2* imprinted locus, a differentially methylated region (DMR) serves as an epigenetically regulated chromatin insulator to bring about the reciprocal imprinting of these two genes (Bell and Felsenfeld 2000). As discussed in the Introduction, regulation occurs through the differential binding activity of CTCF, a DNA binding protein involved in many aspects of gene regulation. CTCF binds the unmethylated maternal *H19/1gf2* DMR, thereby acting as an allele-specific insulator. Putative CTCF binding sites are also found associated with the *D1k1-Gt12* imprinted locus (Takada et al. 2002). To investigate whether CTCF may be involved in the imprinting of the transposed imprinted genes, we first used bioinformatics to predict putative CTCF binding sites. We then verified their *in vivo* binding by PCRbased detection of chromatin-immunoprecipitated DNA (ChIP-PCR), using material from interspecific mouse hybrids to uniquely identify the parental origin of the alleles. We identified two allele-specific CTCF binding sites, one near *Frat3*, the most telomeric imprinted gene in the mouse syntenic region, and a second in the intergenic region between the IC and the transposed imprinted genes. To better understand the epigenetic context of these regions, we investigated histone modifications for the CTCF binding sites as well as their allelic pattern of DNA methylation, and correlated these with the expression patterns of transcripts associated with each site. These results represent the first evidence for CTCF function in the imprinting of PWS region genes.

Results

Identification of potential CTCF binding sites in the mouse PWS/AS region

In order to determine a possible role for CTCF in imprinting of PWS/AS genes, we established criteria for the identification of putative CTCF binding sites from primary DNA sequence, and then identified potential CTCF-binding sites within the PWS/AS region. The CTCF protein can bind to a wide variety of DNA sequences in a methylation dependent manner. A CTCF binding site consensus sequence (CCGCNNGGNGGCAG) was previously derived from the *H19/IGF2* imprinted cluster and the X-chromosome (Chao et al. 2002). We used this sequence as our search query, but also allowed for up to three additional single nucleotide deviations from the core consensus sequence in order to identify a wide distribution of possible CTCF-like binding sites. We searched the DNA sequences published in the February 2003 freeze of the UCSC Genome Bioinformatics database (Karolchik et al. 2003). Specifically, we selected a 2.24 Mb genomic region

defined as 87 kb centromeric of Snurf-Snrpn to 80 kb telomeric of Frat3, which included the transposed imprinted genes Frat3, Mkrn3, Magel2 and Ndn. While the sequence was only available in fragments, the vast majority of these fragments were mapped, oriented, and separated by small gaps. As expected, using low stringency search criteria, we found numerous putative CTCF binding sites, totaling 1558 over 2.24 Mb. We then selected regions containing at least four closely spaced CTCF-like sites clustered within 100 bp of each other, which gave a reasonable number of clusters for our initial analysis. We identified multiple clusters of putative CTCF binding sites in the CpG islands of Ndn, Magel2, Mkrn3 and Frat3 (Figure 5-1). There were also putative CTCF binding site clusters in the CpG islands associated with Snrpn exon 1 (Shemer et al. 1997). The intergenic region between Ndn and Snrpn/IC is rich in retroviral repetitive elements, and these repeats contribute to the almost one Megabase difference in the size of this region between mouse and human (~2.24 Mb vs. ~1.5 Mb respectively). Out of 16 CTCF-like clusters in the intergenic region, there were two clusters (intergenic CTCF clusters A and B, hereafter named iCT-A, iCT-B) that were free of repetitive elements. Cluster iCT-A is a CG rich sequence upstream of a predicted mouse gene annotated with the designation of A230097C02, which is supported by multiple spliced ESTs from embryonic head and hypothalamic libraries. The predicted mRNA encodes a putative protein product of 137 amino acids with no significant similarity to known proteins. There is no human counterpart of the A230097C02 transcript in the homologous region. Finally, as predicted by the CG rich nature of the CTCF-like binding core consensus, we identified many CTCFlike binding site clusters in CpG rich regions, including those associated with transposable

elements. While these elements may be able to bind CTCF *in vivo*, we did not analyze these further as their repetitive nature prevented unambiguous verification of binding using PCR-based techniques.

Figure 5-1. Predicted CTCF binding clusters.

At the top of each figure is a schematic of mouse chromosome 7C, the region of conserved synteny with the human 15q11-q13 PWS imprinted domain. The map is oriented from centromeric (right) to telomeric (left), with a gap between *Snrpn* and A230097C02 of 1.55 Mb. Boxes indicate genes with their name and transcriptional direction above. Diamonds represent CTCF-like binding sequence clusters assayed and their genomic locations. Each figure represents sequence analysis of a section of 7C containing the CTCF clusters of interest. Each vertical line represents one match, with strand and mismatches indicated. CpG dinucleotides are highlighted. (A) region around exon 1 of *Snurf-Snrpn*, (B) iCT-A and iCT-B, (C-D) Region around transposed imprinted genes.







C



Verification of CTCF binding in vivo by chromatin immunoprecipitation

To test whether the predicted CTCF binding sites did indeed bind CTCF *in vivo*, we used PCR-based chromatin immunoprecipitation (ChIP) to assay the region surrounding each of the binding site clusters. The CG rich nature of the cluster at the CpG island starting at position -210 upstream of the start codon of *Frat3* precluded consistent PCR based analysis. Therefore a second sequence upstream of the transcript from -781 to -579 was amplified from ChIP DNA. Sequences for PCR amplification in *Mkrn3* and *Ndn* were also chosen 5' to the putative CTCF binding site clusters (-738 to -552 and -582 to +117, respectively). The repetitive nature of the CpG island of *Magel2* precluded PCR analysis. For *Snurf-Smpn*-IC, a sequence within the CG rich region near exon 1 was analyzed (+60 to +286 with respect to *Snurf* ORF start codon). Regions overlapping predicted CTCF-like clusters were analyzed for iCT-A and iCT-B. The region surrounding the putative CTCF binding site clusters that were chosen for further ChIP analysis were sequenced in DNA from *Mus musculus* (C57BL/6) and *Mus castaneus*. In all cases, we identified polymorphisms that enabled allele-specific analysis in F1 hybrids of crosses between these two species of mice.

ChIP was performed with antibodies against CTCF, using brain and liver tissues obtained from three week old F1 mice generated from a cross between *Mus musculus* (C57BL/6) females and *Mus castaneus* males (Figure 5-2). As a positive control, we tested the DNase I hypersensitive site 1' (HS1') between *TCR* and *Dad1*, which has previously been shown to be highly enriched for CTCF binding compared to a site near HS6 (Magdinier et al. 2004). Both positive control sequences were amplified by PCR from the

DNA obtained by CTCF ChIP, and consistent with published results, we observed a significant difference in CTCF binding between the two sites (Figure 5-2 B). Analogous to findings for the H19/Igf2 DMR (Bell and Felsenfeld 2000), we predicted that the putative CTCF binding site clusters near the PWS/AS IC might bind CTCF preferentially on the unmethylated paternal allele. Contrary to that expectation, there was no allele-specific binding to the CG rich regions associated with the IC, as both alleles displayed relatively little CTCF binding (Figure 5-2 A). In contrast, the region surrounding the cluster of putative CTCF binding clusters associated with Frat3 showed strong, statistically significant binding on the paternal allele versus the maternal allele. This result was found in two independent trials and was observed in both brain and liver samples. Furthermore, in the region surrounding the intergenic cluster iCT-A, we observed binding with a paternal bias (p>0.05 and p>0.1 for the two trials). No significant binding of CTCF to predicted CTCF binding clusters associated with Ndn and Mkrn3 CpG islands was observed on either parental allele. High background levels in the CTCF binding assay at iCT-B precluded quantitative analysis of this site. This is likely due to inefficient probe labeling or annealing, but can also be due to other factors such as probe and primer design.



Figure 5-2. CTCF ChIP in mouse telomeric PWS/AS region.

Quantification of allelic differences in CTCF binding at predicted CTCF-like clusters by chromatin immunoprecipitation. Shown here is representative semi-quantitative CTCF ChIP-PCR data. The y-axis indicates binding compared to input, defined as 1% of starting material. (A) The relative amount of binding to the maternal allele (white) and paternal allele (gray) as measured by hybridization of allele-specific oligonucleotides. Double daggers indicate that allelic differences in both trials gave a t-test P-value of less than 0.05, while daggers indicate one trial gave a P-value less than 0.05 and the other less than 0.1. This is interpreted as an allelic bias. (B) Relative amounts of CTCF binding to the intergenic sites between *TCRa* and *Dad1*, performed as a positive control for the ChIP reaction.

Histone modification at putative CTCF cluster sites

To test whether allelic CTCF binding at Frat3 and iCT-A but not at Ndn, Mkrn3 or the IC was correlated with chromatin context, we used chromatin immunoprecipitation with antibodies that recognize specific histone modifications associated with transcriptional activity. Histone H3 acetylation is typically associated with open active chromatin (Razin 1998). We detected allelic biases in histone acetylation at the Ndn and Snurf-Snrpn CTCFlike clusters, consistent with the expression patterns and imprinted state of these two genes (Ozcelik et al. 1992; MacDonald and Wevrick 1997). The Ndn CTCF cluster had a strong paternal enrichment in histone acetylation (p>0.05, both trials, Figure 5-3 A) in the brain where it is expressed from the paternal allele, but not the liver where both alleles are silent. Expression of the Snurf-Snrpn gene was detected in both brain and liver, and we observed strong paternal enrichment in histone acetylation in the brain (p>0.05, both trials) and a bias towards paternal enrichment in the liver (p>0.05 and p>0.1 for the two trials), consistent with a previous observation (Fournier et al. 2002). We assessed the expression patterns of Frat3, Mkrn3 and the transcript associated with the intergenic cluster iCT-A (A230097C02) in the brain and liver of 3wk old mice by reverse transcription-PCR (RT-PCR) (Figure 5-4). While expression of *Frat3* was undetectable by RT-PCR from these tissues, the region surrounding the Frat3 CTCF-like cluster carried strong paternally enriched histone H3 hyperacetylation in brain (p>0.05, both trials). Mkrn3 expression was detected in the brain and not the liver (Figure 5-4), yet it carried strong paternal enrichment for histone H3 acetylation in both tissues (p>0.05, both trials, both tissues). The intergenic cluster iCT-A showed strong paternal enrichment in histone H3 acetylation in the brain
(p>0.05, both trials), suggesting that the expression of its nearby transcript, A230097C02, in the brain may be paternal allele-specific. Overall, histone acetylation was enriched on the paternal allele of each imprinted gene tested gene, despite the absence of expression of the associated RNA transcript in some cases.

Next, methylation at lysine 4 of histone H3 was examined (Figure 5-3 C-D). Diand tri-methylation of lysine 4 (H3K4me2 and H3K4me3 respectively) is associated with gene activity in different systems (Sims et al. 2003; Schneider et al. 2004). Consistent with a previous report (Fournier et al. 2002), the CTCF cluster near *Snurf-Snrpn* carried an enrichment of H3K4me2 on the paternal allele (p>0.05, both trials, Figure 5-3 C). At the same site, we also observed a bias towards enrichment of H3K4me3 on the paternal allele (p>0.05 and p>0.1 for the two trials, Figure 5-3 D) in both brain and liver. A bias towards paternal allele enrichment in H3K4me2 (p>0.05 and p>0.1 for the two trials) but not H3K4me3 was observed in *Ndn* in brain only. *Mkrn3* carried a bias towards the paternal allele in H3K4me2 in the brain (p>0.05 and p>0.1 for the two trials), while in the liver, H3K4me2 was paternally enriched (p>0.05, both trials) and H3K4me3 was biased towards the paternal allele (p>0.05 and p>0.1 for the two trials), while in the liver, H3K4me2 was paternally enriched (p>0.05, both trials) and H3K4me3 was biased towards the paternal allele (p>0.05 and p>0.1 for the two trials). No significant difference was found for either histone methylation modification in the CTCF binding site cluster near *Frat3*. Finally, the region surrounding the putative CTCF binding sites at iCT-A showed paternal allele enrichment of H3K4me2 and H3K4me3 (p>0.05, both trials, each).

Dimethylation at lysine-79 of H3 (H3K79me2) is associated with developmentally active chromatin regions, and hypomethylation of the same residue has been associated with silencing in yeast (Ng et al. 2003). We found paternal biases in H3K79me2 in the

clusters associated with *Ndn* and iCT-A in the brain (p>0.05 and p>0.1 for the two trials of each, Figure 5-3 B), while *Snurf-Snrpn* carried strong paternal enrichment for H3K79me2 (p>0.05, both trials). We investigated di- and tri-methyl states of the K9 residue, associated with silencing in many systems (Jenuwein and Allis 2001), but found either weak or inconsistent binding to most sites (Figure 5-3 E-F). A maternal bias for tri-methylated H3 K9 was seen in the *Ndn* CpG island (p>0.05 and p>0.1 for the two trials).





Histone modifications of the putative CTCF binding site clusters. Shown here are representative semi-quantitative histone ChIP-PCR data: The data are presented as in (Figure 5-2). (A) Acetylated histone H3, (B) dimethylated lysine 79 of H3, (C) di-methylated lysine 4 of H3, (D) trimethylated lysine 4 of H3, (E) di-methylated lysine 9 of H3, (F) trimethylated lysine 9 of H3.











Figure 5-4. Expression of transposed imprinted genes.

RT-PCR of transposed imprinted genes in 3 week old liver, brain and parts of the brain. For each tissue, the left lane is with reverse transcriptase, while the right lane is without. *Mkrn3* shows different expression patterns in certain parts of the brain. RT-PCR was performed for 30 cycles.

Overall, the CTCF binding cluster iCT-A showed the most striking allelic differences in that it carried paternal enrichment in acetylated H3, H3K4me2, H3K4me3 and a bias in H3K79me2 in the brain, as well as a paternal bias in CTCF binding in both tissues. The paternal allele-specific enrichment and bias of several H3 modifications in spite of the absence of gene expression suggests an allelic differential chromatin structure even in tissues not actively transcribing the transposed imprinted genes. However, no allelic histone H3 modification was perfectly correlated with allelic CTCF binding *in vivo* at any of the tested sites.

DNA methylation patterns of Frat3, Mkrn3 and iCT-A CTCF cluster sites

Binding of CTCF to DNA is prevented by methylation of the target sequence. To test if allelic DNA hypomethylation of the CpG island of *Frat3* is correlated with the allelic pattern of CTCF binding, we performed bisulfite sequencing on DNA extracted from brain and liver of F1 mice. Seventy seven CpG dinucleotides were assayed for methylation. Of these, seventeen are located within ten of the CTCF-like sites in this cluster of thirteen predicted sites (Figure 5-5 A). In both brain and liver, the maternal allele carried moderate methylation (31% and 24% respectively) across all 77 CpG sites, but the paternal allele carried very little methylation (3% and 2% for brain and liver, Figure 5-6 A-D). The CpG dinucleotides within the putative CTCF sites carried 20% and 18% methylation on the maternal allele in brain and liver, and 0.6% and 0% methylation on the paternal allele in brain and liver respectively. Ten CpG dinucleotides were assayed in *Mkrn3* (Figure 5-6 E-H), four of which were located within the putative CTCF binding sites in this cluster of five sites (Figure 5-5 B). The maternal allele of *Mkrn3* in the brain carried much higher levels

of methylation (83%) than the paternal allele (7%) in the brain; equivalent methylation in liver was detected for the two alleles. Methylation levels of the CpG dinucleotides within putative CTCF binding sites were found to be similarly high on the maternal allele in brain and both alleles in liver (82%, 78%, and 58% respectively) but low on the paternal allele in brain (8%). Eight CpG dinucleotides were assayed in the iCT-A intergenic cluster (Figure 5-6 I-L), two of which are within four putative CTCF binding sites in this cluster of seven sites (Figure 5-5 C). As in the sites near Frat3, DNA methylation of iCT-A was found to be higher on the maternal allele in both brain and liver (55% and 45% respectively) than the paternal allele (0% and 13% in brain and liver). The putative CTCF binding sites on the paternal allele of Mkrn3 are hypomethylated in the brain but were not found to bind CTCF (Figure 5-6 F), and thus DNA hypomethylation is not sufficient for CTCF binding to these sites. In this case, the CpG dinucleotides within putative CTCF sites had similar levels of methylation to those not in CTCF sites. Overall, all alleles that showed in vivo CTCF binding also displayed hypomethylation of the CpG dinucleotides located within the predicted CTCF binding sites on the paternal allele, as well as hypomethylation of sites in the rest of the CpG island, when compared to the maternal allele.

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	88661	TOOOGACAGO	TOTOCATACC	CCCATGGAGA	ANTATO <u>CG</u> AC	ATGTGCOGGC			
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P(A/GCCAOT)

06

B

Figure 5-5. CpG sites analyzed for DNA methylation.

Regions analyzed by bioinformatics search for CTCF-like binding sites, which are highlighted red and green (upper and bottom strand, respectively), and by bisulfite sequencing. CpG dinucleotides are underlined and numbered. Polymorphisms are indicated by a P, followed by the *M. musculus* and *M. castaneus* variants. Downward arrowhead in (A) indicates 5' end of region deleted in a Frat3 gene targeted mouse strain.(A) *Frat3*, (B) *Mkrn3* and (C) iCT-A.



Figure 5-6. DNA methylation of Frat3, Mkrn3 and iCT-A.

DNA methylation patterns on the maternal and paternal alleles of *Frat3*, *Mkrn3* and iCT-A. The y-axis indicates the percentage of clones that are methylated at each CpG dinucleotide site marked on the x-axis. The bisulfite-treated DNA samples were isolated from brain and liver tissue samples. Sites are numbered as in Figure 1. Black and gray bars indicate CpG dinucleotides within or outside of predicted CTCF binding sites respectively (A-D) *Frat3*. (E-H) *Mkrn3*. (I-L) iCT-A.

Discussion

CTCF binds its target sequence through combinatorial use of eleven zinc finger domains (Klenova et al. 2002). The variation in the use of these domains leads to a considerable variation of CTCF in vivo DNA binding sites. The use of specific zinc fingers to bind subsets of CTCF targets also restricts which zinc fingers remain available for interactions with other proteins involved in CTCF functions. Thus, we reasoned that the consensus CTCF binding sequences found at loci where CTCF was involved in monoallelic gene regulation, such as X-inactivation and imprinting of H19/IGF2 (Chao et al. 2002), could be shared with CTCF functions at the PWS/AS locus. Using this strategy, we searched the available mouse DNA sequence that has conserved synteny with the human PWS/AS region for a CTCF binding site consensus sequence. This strategy has previously proved useful in finding novel CTCF binding sites at the H19/Igf2 locus (Ishihara and Sasaki 2002). As this study was designed to test the most likely CTCF binding sites, others may well exist in the PWS/AS region, and this awaits more comprehensive approaches such as those on genomic microarrays. Out of 21 predicted clusters of CTCF binding sites, five were amenable to ChIP-PCR analysis, and we found that only two bound CTCF in vivo. Of great interest was the observation that both CTCF binding clusters showed paternal allele-specific CTCF binding. Notably, we also observed that the DNA of the paternal allele was hypomethylated in both tissues studied, correlating with allelic CTCF binding.

Allelic histone modifications and DNA methylation have previously been shown to be associated with imprinted genes (Delaval and Feil 2004). DMRs are associated with 141 many imprinted genes and some carry germline imprints. Likewise, several histone modifications are found with many imprinted genes (Fournier et al. 2002). Our data suggest no single epigenetic modification is synonymous with imprinted gene regulation and that differences exist between species and even genes in the same cluster. Our previous studies of human NDN showed that methylation of H3 lysine 4 was well correlated with allelic identity while histone acetylation was correlated with the imprinted and tissue-specific expression of NDN (Lau et al. 2004). In this study, paternal allele-specific H3 lysine 4 methylation and histone acetylation of Ndn were both correlated with expression in the brain, while no allelic histone modifications were found in the liver, where Ndn is not expressed. Species-specific differences in DNA methylation patterns were also observed. DNA hypermethylation was associated with the maternal allele in human NDN regardless of expression (Lau et al. 2004). In the mouse, similar allelic differences were found in the brain, where *Ndn* is expressed in many cells, but both alleles were hypomethylated in the heart, where Ndn is not expressed (Hanel and Wevrick 2001). This may be due to selection of representative expressing and non-expressing tissues, since fibroblast and lymphocytes were studied in human and brain and heart were studied in mouse, respectively. There may be either species or tissue-specific differences in the relationship between epigenetic modifications and allele identity. In our current comparison of the transposed imprinted genes in the same cluster as Ndn, no definitive epigenetic signature was shared by this cluster of genes. While all genes tested carry allelic histone modifications to some degree in at least one tissue, these modifications do not correlate well with the expression pattern of each individual gene. This may be due to the regions chosen for analysis, since our work

(Lau et al. 2004) and work by others has shown that histone modifications can have very restricted distribution patterns (Schneider et al. 2004). We did observe a general theme of active modifications such as DNA hypomethylation, histone acetylation, and histone methylation on lysine 4 and 79 of H3 on the paternal allele even in some cases where there was absence of expression. This is consistent with a model whereby the paternal allele is more open and carries active modifications, although the repression of transcription depends on tissue-specific factors that act in spite of the permissive modifications at the locus.

Since *Frat3* is not expressed in the adult tissues we tested, it is unlikely that the differential CTCF binding affects transcription, or that the DNA methylation or histone acetylation pattern is related to transcription. Given the critical role CTCF plays in *H19/Igf2* imprinting (Bell and Felsenfeld 2000), and the mechanistic model by which CTCF functions in allele-specific higher-order chromatin structure formation, we propose that the CTCF clusters that we identified also operate to organize chromatin in an allele-specific manner. The germline-derived differentially methylated region (DMR) of *H19* is unmethylated on the maternal allele, binds CTCF, and forms a complex with the upstream maternally unmethylated DMR1 of *Igf2*. This complex forms a loop bringing *Igf2* into an inactive nuclear domain. On the methylated paternal allele of the *H19* DMR, an unknown protein complex is instead formed with the paternally methylated DMR2, at the 3' end of *Igf2*, bringing *Igf2* out of the loop and theoretically giving it access to a nuclear environment permissible to transcription (Murrell et al. 2004). Countless configurations of the PWS region are possible, although one model analogous to the *H19/Igf2* paradigm can

be postulated. The *Frat3* CTCF site and the iCT-A site may cooperate to fill the role of a matrix anchor (Yusufzai and Felsenfeld 2004) and bring the transposed genes as a coregulated unit in and out of active nuclear regions on the paternal and maternal alleles respectively in response to the IC (Figure 5-7 A and B). The *gypsy* insulator of *D*. *melanogaster* also functions through changes in nuclear localization (Gerasimova et al. 2000). An overall differential chromatin context for the co-regulated unit would also be consistent with our observations that some of the histone modifications we examined were allele-specific without concurrent expression of the associated transcript, in particular *Frat3* itself. It also remains to be seen whether the allelic CTCF binding is conserved in humans as is the case with the *H19/Igf2* gene pair.

Figure 5-7. Model of CTCF function at transposed imprinted genes.

Model for the function of CTCF binding in the transposed genes of PWS/AS 7C imprinted domain. (A) On the paternal allele, flanking CTCF binding sites (filled circles) are bound by CTCF (half donuts) and the intervening transposed genes are sequestered into nuclear compartments permissible to gene activity, thus allowing a euchromatic state (single line). The expression of intervening genes and the state of histone modifications and DNA methylation are dependent on species- and tissue-specific factors. (B) On the maternal allele, the CTCF binding sites are not bound by CTCF, allowing heterochromatinization of the transposed genes along with the rest of the imprinted domain on that allele. (C) Upon deletion of the telomeric CTCF binding site of *Frat3* (empty circle), the paternal allele may partially lose association with active nuclear compartments, allowing position effects of chromatin outside the domain (double line) to encroach. The degree of this effect remains to be tested in this deletion mouse.



Whether the CTCF binding sites identified in this study participate in intralocus association within the PWS region in a manner analogous to the H19/Igf2 locus await further study, with these two sites being ideal starting points for the chromatin conformation capture assay (Dekker et al. 2002). The CTCF binding site at Frat3 may have an alternate role in demarcating the telomeric end of the imprinted domain at mouse 7C and may insulate the domain from more telomeric genes, or conversely, may protect other genes from the effects of the long range actions of the IC. Targeted deletion of the CTCF binding sites associated with Frat3 and the iCT-A clusters would test the hypothesis that these are indeed important in the maintenance of the imprinted states for all four transposed imprinted genes. Three independent gene targeting experiments that modified Ndn, Frat3, and the highly related but non-imprinted *Frat1* gene are useful in elucidating the possible importance of the distal CTCF binding site cluster in imprinting. In one Ndn gene-targeted line, 33% of the CpG island and most of the open reading frame of Ndn was replaced with LacZ, which became imprinted under control of the Ndn promoter (Gerard et al. 1999). A gene-targeted deletion of Frat3 that includes most of our predicted CTCF-like sites was recently reported (van Amerongen et al. 2005). In this case, the Frat3 open reading frame, which contains most of the CTCF-like sites, was replaced with a LacZ reporter gene. This removed 80% of the CpG island and ten of thirteen CTCF-like binding sites. Surprisingly, the reporter became silent and methylated on both alleles. Frat1 is the highly homologous ancestral gene to Frat3, and is located in a separate region of the genome and is therefore not imprinted. Gene-targeting of Frat1 removed its open reading frame, coincident with 77% of the CpG island, and allowed correct spatial-temporal embryonic expression of the

inserted reporter gene (Jonkers et al. 1999). While the silencing of *Frat3* could be due to removal of basal promoter elements of *Frat3*, the fact that a very similar *Frat1* deletion still enabled correct expression of a reporter gene argues against this hypothesis. We favor the interpretation that the *Frat3* deletion removed CTCF binding sites critical for activity on the paternal allele, which normally protects *Frat3* from position effects telomeric to the imprinted domain (Figure 5-7 C). A similar role for CTCF sites has been proposed for the delimitation of the imprinted *Igf2/H19* and non-imprinted β -globin domains (Saitoh et al. 2000; Ishihara and Sasaki 2002). Functional insulator assays and matrix enrichment assays of this CTCF binding site will help to refine this model. Further examination of expression and epigenetic marks of the other three transposed genes in the *Frat3* transgenic mouse model could be instrumental in understanding the role of the *Frat3* CTCF binding site in genomic imprinting.

The possible role of CTCF in imprinted regulation and maintenance of the PWS domain has been poorly defined. Discovery of the first evidence of CTCF involvement within the PWS imprinted region was made possible by our strategy based on clustering of multiple CTCF-like consensus binding sites. This serves as a critical first step in a more comprehensive documentation of CTCF binding and function in imprinting of PWS genes as well as other large imprinted clusters. Our results here also raise the possibility that the transposed imprinted genes may be regulated through a mechanism similar to the *H19/Igf2* locus, while the genes centromeric (in the mouse) to the PWS IC are imprinted *via* a mechanism involving transcription of an antisense RNA (Chamberlain and Brannan 2001; Runte et al. 2001) in a manner similar to that proposed for the *Igf2r/Air* locus (Sleutels et

al. 2002; Landers et al. 2004) (Figure 5-8). The mechanisms whereby the PWS/AS IC can bring about imprinting by more than one mechanism merit further study. With detailed characterization, this unique feature of the PWS/AS imprinted cluster may begin to reconcile seemingly disparate models of imprinted regulation at different loci.



Figure 5-8. Models of imprinting operating at PWS/AS domain.

Comparison of different models of imprinted regulation. The interpretation of the data presented here raises the possibility that while the centromeric side of the PWS cluster may employ the antisense suppression mechanism similar to X-inactivation and other imprinted clusters such as the Igf2r/Air locus, the telomeric side may make use of allelically regulated CTCF binding similar to the Igf2/H19 locus.

Chapter 6 Conclusions and Open Questions

The phenomenon of genomic imprinting has enjoyed the attention of many investigators ever since its discovery. By breaking from Mendel's rules, imprinting has become one of the last great mysteries in the study of human heredity. As well as being an irresistible biological puzzle, it also has relevance to human congenital disease, development, oncogenesis and evolution. Imprinting also shares many aspects of gene regulation with other systems such as X-inactivation and regulation of the well studied β globin cluster. Therefore, detailed investigation of genomic imprinting will give insight into mechanisms applicable to general gene regulation, and will better define the role of imprinting in the scheme of human biology.

The PWS/AS cluster is an excellent example of an imprinted locus. There are well defined human disorders based on isolated deficiency of both alleles, and mutations defining the imprinting center that carries the germline imprint. A well conserved region of synteny in the mouse presents a convenient model organism for genetic dissection and evolutionary conservation. Evidence suggests genes centromeric and telomeric to the IC are controlled by different mechanisms. While a partial model exists for the mechanism of imprinting for one side of the locus (telomeric in human, centromeric in mouse), little is known about the regulation of the transposed imprinted genes. A well-defined imprinted gene, *NDN/Ndn*, was used to investigate the less defined aspects of imprinting at this locus. By studying this gene from varying perspectives, starting with fine scale examination of its promoter, then histone composition for a region encompassing the gene, to domain wide regulation, these studies have given surprising insight into imprinting of the transposed imprinted genes in the PWS region.

Regulation of the transposed imprinted genes

Detailed bioinformatic examination of the promoter sequence of *NDN/Ndn* from human, mouse and rat has shown conservation of promoter sequences, as well as putative binding sites for several transcription factors. Certain features such as conservation in sequence of part of the minimal promoter and predicted sites for factors involved in the control of genes in neuronal function are consistent with some proposed roles for *NDN/Ndn*, since there are overlapping expression patterns of human and mouse in the nervous system. Comparison of this phylogenetic footprint to the *in vivo* footprint information gathered by Dr. Meredith Hanel has been useful in suggesting avenues of investigation on the identity of the *trans*-acting factors regulating *NDN*. While the candidate proteins tested here have not been proven to regulate *NDN* at its promoter, other candidates identified in these studies may prove themselves functional in the future. More importantly, the data presented in Chapter 3 has suggested chromatin accessibility, instead of a purely *trans*-factor based scheme, is also involved in the tissue and allelic regulation of *NDN*.

To further characterize differences between parental alleles and expressing and non-expressing alleles, the chromatin context in terms of histone modifications was defined for *NDN* in Chapter 4. As expected, certain modifications known to be associated with activity, such as histone acetylation and methylation at lysine 4 of H3, are associated with the paternal allele in an expressing cell type. In this instance, the modifications also displayed a distinctive distribution where acetylation was spread out over a wide area including sequences upstream and downstream, while dimethylation of lysine 4 was more

restricted and trimethylation was only found in a small region within the region of dimethylation. Surprisingly, methylation of lysine 4 of H3 was also found to be paternal allele-specific in the non-expressing cell type tested, although with a less distinct distribution. Therefore, there seem to be different types of "active" chromatin at NDN. The paternal allele can carry the active modification of lysine 4 methylation and DNA hypomethylation, whereas in a cell type where NDN is also transcribed, acetylation also occurs. This may indicate that lysine 4 methylation is a persistent allelic mark regardless of expression. This is correlated to the situation in the *in vivo* footprint, where the transcription-related footprint is associated with the paternal allele in the expressing cell type, but other footprints are allelically associated in both cell types. Of course the cause and effect relationship between binding of protein factors and histone modifications cannot be determined in these assays. The hierarchical dominance of one over the other is also not mutually exclusive. One scenario can be envisioned where histone lysine 4 methylation represents a persistent mark of the paternal allele, transmitted through cell divisions, and may allow binding of certain factors, some in every cell, and some in cells expressing NDN. The binding of factors up-regulating NDN in expressing cells may recruit acetylases and chromatin remodelers to aid in opening up the locus and allow binding of other factors and transcriptional machinery. Identification of the factors involved in regulating NDN will help define this relationship. Further characterization of histone modifications in other tissue types and developmental stages will also be useful in determining the order of events in the regulation of NDN.

To gain a better understanding of the mechanism involved in the differential chromatin states of the two alleles, it was necessary to take a step back and look at the big picture. Since CTCF had previously been shown to have a role in imprinting and long distance gene regulation (Kato and Sasaki 2005), its involvement in the imprinting and regulation of the transposed imprinted genes in the PWS/AS cluster was tested. A bioinformatics search for binding sites similar to previously characterized sites revealed candidate regions where CTCF was predicted to bind. Verification of binding *in vivo* made possible a novel discovery of a possible regulation model for these genes. Two sites flanking the transposed imprinted genes were found to bind CTCF, and did so only on the paternal allele. These sites may delimit a co-regulated domain containing this set of genes and aid in their imprinted regulation. Again, cause and effect cannot be determined at this point, and whether CTCF causes allele-specific organization or differential chromatin states allows CTCF binding awaits further study.

Since the genetic evidence in human and mouse suggests no locus other than the IC carries the germline imprint, it is likely the CTCF sites identified here are bound in response to the IC on the paternal allele. One of the ways in which this can be controlled is through differential DNA methylation (Figure 6-1). Positive influence of the IC on the paternal allele may exclude DNA methylation of CTCF sites and allow binding. Alternatively, heterochromatinization of the maternal allele may recruit DNA methylating activities to these sites. The binding of the flanking CTCF sites may lead to sequestration of the domain into a permissive nuclear address where it could be accessible to modifying enzymes, marking the allele with euchromatic histone modifications. These modifications

and differential DNA methylation at the CTCF binding sites may contribute to the somatic memory of the paternal allele throughout development. When the paternal allele arrives in a tissue that carries factors necessary in the transcriptional activation of one of the transposed imprinted genes, they are able to bind the paternal allele at the permissive nuclear address, and affect further accumulation of active histone modifications, leading to opening of the chromatin structure and gene activation. This model fills a void in the understanding of the PWS/AS imprinted domain. As described in the Introduction, the antisense model of imprinting satisfies the observations for many of the imprinted genes such as *SNURF-SNRPN*, snoRNAs and *UBE3A* (Runte et al. 2001). The allelic binding of CTCF at the transposed imprinted genes complements the antisense model with a mechanism to address imprinting of these genes where the antisense mechanism is insufficient.

Figure 6-1. Model of epigenetic relationship between imprinted and tissue-specific

regulation.

Schematic of a model of the relationship between chromatin accessibility, histone modifications and CTCF binding to the transposed imprinted genes. An initial signal from the IC in *cis* determines differential epigenetic states on the two alleles, including allelic DNA methylation (lollipop) at CTCF sites (dark green circles) flanking transposed imprinted genes (pentagonal block arrow). This then leads to allelic binding of CTCF (dark blue half donuts), and association of the paternal allele to active regions of the nucleus (green haze), with access to factors that can reinforce a differential epigenetic state. The other allele would lack CTCF binding and be associated with a different set of factors (grey octagons). Tissues expressing one of the transposed imprinted genes would contain tissue-specific factors (orange triangles) that will have access to the paternal allele, and further add active epigenetic marks such as histone modifications and lead to transcription of the paternal allele in that tissue (green pentagonal block arrow).



Early events of the transposed imprinted genes

As with all new discoveries in science, answers spawn more questions, and the work presented here will lead to future efforts in filling in details of this model. The importance of determining the order and hierarchy of regulatory events is evident from this work. The experiments described here address the state of imprinting of the transposed imprinted genes in somatic tissues. However, the allelic differences observed were likely not set up *de novo*, but epigenetically inherited. There are two distinct questions here: 1) the question of the order of events early in the setting up and interpretation of the original imprint, and 2) the setup and propagation of the imprint that survives cell divisions in the soma. As was discussed in the Introduction, the order is not even clear at the IC, nor is a complete picture available for any imprinted domain.

Allelic imprints of the previous generation are erased during germline formation through an active global demethylation event (Figure 6-2) (Hajkova et al. 2002). This complete erasure allows each generation to mark imprinted genes according to gender. During gametogenesis, the exclusive expression of the CTCF-related protein, BORIS in the male germline has been postulated to be important in preventing methylation of imprinted ICs that would otherwise be methylated in female gametogenesis (Loukinov et al. 2002). The female germline however, without the need to endure the dramatic genome-wide repackaging required by spermatogenesis, may use DNA methylation, histone modifications or binding of other protein factors to dictate imprinted states. The methylation state of the PWS/AS IC is likely to fit into these schemes and survive the subsequent zygotic demethylation with the maternal IC methylated and paternal IC

unmethylated. While DNA methylation for ICs is known to be allelic at this stage, there may be other epigenetic marks present, such as histone modifications. CTCF, the relatively new player on the block, may be involved in the early events in X-inactivation (Pugacheva et al. 2005), but its significance at the PWS/AS IC has not been determined. In general, the hierarchy of control has plagued the field of gene regulation. Instead of a linear hierarchy, there may be a network of overlapping epigenetic signals reinforcing and fine tuning the regulation of a gene. This view may be a compromise but still does not suggest a candidate initiating signal driving allelic expression of the transposed imprinted genes. Detailed description of the correlation between other imprinted genes and between different species in terms of the language of epigenetic modifications will also help our understanding of the importance of different marks over others. Undoubtedly, whole genome approaches will be instrumental in this debate (Bernstein et al. 2005).

Figure 6-2. Cycle of imprinted inheritance of the IC.

Establishment of the imprinted state as observed in somatic tissues (top right) begins in development of the gametes in the previous generation (left). Blue and red lines represent DNA of the paternal and maternal allele, respectively. Differential DNA methylation patterns at ICs are the result of events specific to spermatogenesis and oogenesis. In spermatogenesis, testis-specific factors such as BORIS may play a role in determining DNA methylation states, whereas in the ovary, factors other than DNA methylation may also be employed such as histone modifications or binding of unknown proteins (green diamond). After fertilization and the mixture of the two parental contributions, these epigenetic states must then be elaborated into other epigenetic signals that aid in transmission and transcriptional control of the two alleles. Cells of the soma in the new generation must re-enter this cycle of imprint inheritance to maintain correct genderspecific marks for the next generation.



The allelic identity at the IC must also traverse the genomic distance necessary to reach the imprinted genes in the PWS/AS region. Evidence in PWS patients with somatic deletions of the IC indicates that this is an ongoing process that requires influence of the IC throughout development, instead of an isolated early post-fertilization event (Bielinska et al. 2000). While our studies in somatic tissues only indirectly addressed the initial events determining allelic identity, somatic tissues are ideal for studying the constant requirement of the transposed imprinted genes for an LCR-like function for the IC in imprinting.

Talking to the transposed imprinted genes

Our question of hierarchy is intimately linked to another outstanding question of wide interest in imprinting and gene regulation, and that is how regulatory signals traverse significant genomic distances. Mechanisms involved in both forms of regulation may be one and the same. One of the simplest and oldest models of long range control is physical spreading of epigenetic state. The patterns of DNA methylation and histone methylation that are thought to function in transcriptional regulation may also play the role of long distance communication. This can be achieved through reinforcement of the epigenetic network as described above and in the Introduction (Figure 1-11). For example, H3K9me3 is able to bind the heterochromatic protein HP1, which in turn is able to recruit the H3K9 HMT SUV39H1, which then reinforces the positive feedback loop by methylating H3K9 (Figure 6-3 A) (Grewal and Moazed 2003). This loop is not the only mechanism however, as H3K9me can cause histone deacetylation and silencing independently of HP1 (Stewart et al. 2005).

Figure 6-3. Models of distant epigenetic influence.

Depicted here are several non-mutually exclusive models of distant control by an LCR or IC element. (A) Model of linear spreading of epigenetic modifications, with propagation of H3K9me schematically shown. The epigenetic loop of H3K9me binding of the HMT SUV39H1, subsequent methylation of H3K9 and further recruitment of HMT can feedback and amplify the spreading of the heterochromatic state on the maternal allele from the IC. (B) Model of direct communication between the IC and the transposed imprinted gene cluster (green and red pentagonal arrows) by intralocus association. A protein of unknown identity (orange cross) may bind the IC and form an active chromatin hub (ACH) with CTCF (blue half donuts) bound at a distance. (C) Model of physical or temporal separation of alleles by action of the IC. The IC may cause allelic sequestration of the alleles into different nuclear addresses or differential replication timing. Separation would result in differential access to gradients of activating complexes (green triangles), and silencing complexes (grey octagons).



Looping mechanisms such as described for Igf2/H19 are one of the most common models for long distance control (Bulger and Groudine 1999). Although our data do not support CTCF binding at the IC, the data do not preclude intralocus association between the IC and the CTCF sites flanking the transposed imprinted genes. In fact, it is clear that factors other than CTCF must be able to participate in the loop formation at Igf2/H19 since CTCF only binds the unmethylated DMR, but the methylated DMR, and other sequences found to associate with the DMR in cis do not bind CTCF (Murrell et al. 2004). The identity of these DNA binding proteins are unknown, but it is possible that similar factors can bind the PWS IC in a similar way and mediate association with the CTCF sites flanking the transposed imprinted genes, thus forming an allele-specific active chromatin hub (Figure 6-3 C). It is surprising that no reports detailing the intralocus association of the PWS/AS or any other imprinted domain has been published thus far. However, lack of supporting evidence does not equate to evidence to the contrary, and may simply be due to the technical difficulties involved in assaying a region as large as the PWS/AS domain with present techniques. Intralocus association would certainly explain why the IC is necessary for continued imprinting of the region in somatic tissues. Further characterization of these associations and factors involved will be critical in understanding imprinting of the PWS/AS cluster and how this relates to imprinting in general.

Another perspective in viewing the question of how the IC imparts its allelic signal is to look at the physical separation of the alleles. Initial erasure and establishment of parental imprints occurs in gametogenesis when parental alleles are isolated. Even after
fertilization, the parental genomes remain separate in a stage when germline imprints may be translated to other allelic epigenetic marks. After these stages, it may be a challenge to physically separate the alleles to facilitate differential regulation. A model of sequestration of alleles based on CTCF binding has already been presented here. If the IC functions in a similar way, by allelically determining structural nuclear association, as suggested by others (Greally et al. 1999; Kagotani et al. 2002), this would be another mechanism consistent with a somatic requirement of the IC. An alternative involves temporal separation. Replication timing has been associated with imprinted domains but has not been well characterized in terms of its function in imprinting (Bickmore and Carothers 1995). This represents a unique mechanism of separating alleles. Perhaps histone modifying and chromatin remodeling activities vary during the cell cycle and alleles replicating at different times may gain or only be able to maintain certain states. While the replication timing for the IC has been studied, replication of the transposed imprinted genes could be under a different origin of replication and its timing may be dependent on other factors. Physical or temporal separation of alleles would also represent a solution to long range communication between the IC and the transposed imprinted genes; by simple indirect influence. Communication may not be necessary if the IC can direct the two alleles to completely different nuclear addresses and limit the potential chromatin modifying and transcriptional machinery available. All of these questions will fuel the unfolding of this biological puzzle for years to come.

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