Chromatin modifications associated with diabetes

Short title; chromatin, epigenetics and diabetes

Samuel T. Keating (1) and Assam El-Osta(1,2,3,4)

Corresponding author: assam.el-osta@bakeridi.edu.au
Telephone 613-8532-1389
Fax 613-8532-1100

Word count: 5959

1) Epigenetics in Human Health and Disease Laboratory, 2) Epigenomics Profiling Facility, Baker IDI Heart and Diabetes Institute, The Alfred Medical Research and Education Precinct, Melbourne, Victoria 3004, Australia;
3) Department of Pathology, The University of Melbourne, Victoria 3010, Australia;
4) Faculty of Medicine, Monash University, Victoria 3800, Australia
Abstract

Accelerated rates of vascular complications are associated with diabetes mellitus. Environmental factors including hyperglycemia contribute to the progression of diabetic complications. Epidemiological and experimental animal studies identified poor glycemic control as a major contributor to the development of complications. These studies suggest that early exposure to hyperglycemia can instigate the development of complications that present later in the progression of the disease, despite improved glycemic control. Recent experiments reveal a striking commonality associated with gene-activating hyperglycemic events and chromatin modification. The best characterized to date are associated with the chemical changes of amino-terminal tails of histone H3. Enzymes that write specified histone tail modifications are not well understood in models of hyperglycemia and metabolic memory as well as human diabetes. The best-characterized enzyme is the lysine specific Set7 methyltransferase. The contribution of Set7 to the etiology of diabetic complications may extend to other transcriptional events through methylation of non-histone substrates.
**Introduction**

Diabetes mellitus is characterised by chronic hyperglycemia induced by loss of insulin producing pancreatic β-cells (Type I diabetes) or progressive loss of insulin sensitivity and β-cell dysfunction (Type II diabetes). It is estimated there are more than 1.7 million sufferers in Australia [1] with worldwide estimates suggesting 150 million patients as of 2003. This is expected to increase to 300 million individuals with the disease by 2025 [2]. Both forms of the disease are associated with accelerated rates of microvascular complications including retinopathy, neuropathy, and nephropathy as well as macrovascular complications such as hypertension, atherosclerosis and stroke. Hyperglycemia presents as a major risk factor for the development of diabetic complications, particularly endothelial dysfunction associated with vascular complications [3]. The vascular damage arising from hyperglycemic insult acts upstream of the overproduction of reactive oxygen species (ROS) by the mitochondrial electron transport chain [4]. Importantly this has been demonstrated to activate the NFκB pathway leading to inflammatory events in the vasculature [4].

Several large-scale studies highlight the clinical benefits of strict glycemic control and revealed that early hyperglycemic events instigate the development of diabetic complications that present much later in the progression of the disease [5,6]. In conjunction with studies in experimental animal models and cell culture, these observations imply that a cell may retain a memory of past hyperglycemic events that culminate in altered and persistent gene expression that drives the disease phenotype. This concept has been termed ‘metabolic memory’ or the ‘legacy effect’ [5,6]. Current knowledge suggests that chromatin modifications and epigenetic determinants play a key role in the establishment and progression of metabolic memory and persistent gene expression.

In this review we highlight the epidemiological evidence and experimental studies that demonstrate the detrimental effects of hyperglycemia and support the concept of metabolic memory with a particular focus on Type I diabetes. Furthermore we discuss the current knowledge surrounding epigenetic mechanisms that link hyperglycemia-induced vascular injury to chromatin modifications and altered gene expression as well as metabolic memory. We discuss emerging experimental evidence of changes in genomic methylation and posttranslational modifications such as acetylation and methylation of histone tails, before focusing on perhaps the best characterized methyl-writing enzyme involved in lysine mono-
methylation. Additionally we review recent advances in the understanding of Set7-mediated molecular mechanisms that could drive altered gene expression in the diabetic setting.

**Epidemiological studies**

Large-scale studies have highlighted the requirement for strict glycemic control in delaying the onset of diabetic vascular complications. The Diabetes Control and Complications Trial (DCCT) was conducted with a cohort of 1441 Type I diabetic individuals and a mean follow-up of 6.5 years with the aim to compare conventional and intensive glycemic control regimens [7]. Original findings of the DCCT reported that intensive glycemic control reduced the onset and progression of long-term diabetic complications that included neuropathy, nephropathy and retinopathy compared with patients that received conventional therapy [5]. At the conclusion of this trial, a follow-up observational study, the Epidemiology of Diabetes Intervention and Complications (EDIC) examined the long-term effects of the original DCCT cohort. While the DCCT concluded that the development of cardiovascular complications were indistinguishable across intensive and conventionally treated groups, the EDIC trial identified a significantly lower risk of retinal and renal disease in the group that received intensive treatment [8]. At the conclusion of the DCCT, glycosylated hemoglobin (HbA1c) levels differed by approximately 2% between the two groups [5]. Intriguingly towards the end of the EDIC study, HbA1c levels of both groups had converged to comparable levels [8]. Despite the normalization of HbA1c, the effects of intensive therapy conducted for 6.5 years during the DCCT were persistently beneficial for at least 10 years with regard to microvascular complications. More recently it was reported that patients that received continuous intensive treatment throughout the trials were at significantly lower risk of macrovascular complications including cardiovascular disease and stroke [9], as well as atherosclerosis [10,11]. Long-term benefits of glycemic control have been demonstrated to extend to Type II diabetes mellitus. The UK Prospective Diabetes Study (UKPDS) that compared conventional dietary glucose control with intensive insulin therapy over 10 years initially reported decreased microvascular complications in the intensive therapy group [6]. In accordance with the DCCT-EDIC study, 10-year post-trial follow-up revealed lower incidence of macrovascular complications in the intensive therapy group compared to the conventional treatment group [12]. Overall, such findings indicate that previous episodes of poor glycemic control initiate the deleterious effects on the vasculature that persist despite intensive treatment and normalization of blood glucose.
Metabolic memory

In vivo evidence from animal models

From observations arising from these clinical trials the term ‘metabolic memory’ was introduced to describe the clinical features of diabetic complications that continue to develop long after exposure to the hyperglycemic insult, despite improved glycemic control [5,8,13]. Observations from pioneering studies in animal models also provide strong evidence of this phenomenon. Early studies in dogs with alloxan-induced diabetes demonstrated persistent retinopathy despite a period of 2.5 years of insulin therapy and glycemic control [14]. Similarly sucrose-fed diabetic rats that received islet transplantation after 6 weeks of diabetes exhibited reduced progression of retinopathy compared with islet transplantation following 12 weeks of diabetes [15]. Another study in rats with streptozotocin-induced diabetes showed that retinal oxidative and nitrative stress resulted from periods of hyperglycemia, and that late reinstitution of strict glycemic control was insufficient to inhibit the progression of diabetic retinopathy [16]. Initiation of good glycemic control after 2 months of hyperglycemic conditions had only partial benefits with regard to oxidative and nitrative stress, whereas a delay in this intervention for 6 months resulted in complete failure to reverse any abnormalities [16]. These observations highlight difficulties in reversing hyperglycemia-induced changes and the need for strict blood glucose control early in the progression of diabetes. A study in mice cells revealed novel insights into the gene transcriptional changes that underlie the phenomenon of metabolic memory with particular regard to inflammation. Aortic endothelial cells from mice subjected to a hyperglycemic clamp for 3h exhibited increased expression of the pro-inflammatory NFκB p65 subunit. This increase in gene expression persisted for 6 days beyond normalization of blood glucose [17]. The p65-NFκB transcription factor drives expression of several pro-inflammatory genes that contribute to the development of diabetic vascular disease [18]. Furthermore p65 expression is increased in diabetic ApoE-null mice aorta and appears to be a central mediator of inflammation in diabetic complications [19].

Cell culture experiments demonstrating persistence

Transcriptional changes that govern the pathogenesis of diabetic complications in response to the hyperglycemic insult have begun to be addressed in experiments performed in cell culture. These studies have primarily focused on the vascular endothelium, a key site of hyperglycemia-induced injury [20]. Persistent gene expression was addressed in an early study of cultured primary human endothelial cells exposed to transient high glucose (HG)
concentrations. Hyperglycemia induced expression of fibronectin and collagen IV that was maintained despite a return to normoglycemia [21]. Similarly, transient HG increased expression of p65 in primary endothelial cells. Consistent with observations in experimental animal models, the increase in transcription was maintained for up to 6 days following a return to physiological glycemic conditions [17]. Furthermore, expression of downstream p65-NFκB transactivation targets relevant to diabetic vascular injury, monocyte chemoattractant protein 1 (MCP-1) and vascular cell adhesion molecule 1 (VCAM-1) were significantly increased and remained elevated upon return to normoglycemia [17]. The biochemical basis for the transcriptional response to HG has been delineated by key experiments that implicate the generation of reactive oxygen species (ROS) in the development and persistence of diabetic complications [22]. Increased expression of p65 and downstream transactivation targets was abolished by over-expression of either manganese superoxide dismutase (MnSOD) or uncoupling protein 1 (UCP-1) [17], both of which prevent hyperglycemia-induced superoxide accumulation [4,23]. Similarly the expression of protein markers of hyperglycemia such as protein kinase C-β (PKCβ) and neutrophil cytosol factor 1 (p47phox) subunit of NADPH oxidase in endothelial and retinal cells was normalized by inhibition of ROS [24]. Thus the persistence of transcriptional changes following periods of hyperglycemia seems to be imparted by mitochondrial ROS overproduction. Linking these biochemical changes with nuclear mechanisms that govern transcriptional events is a key objective towards understanding glucose-induced transcriptional changes and metabolic memory. Several recent observations have sparked considerable interest in epigenetic mechanisms of gene regulation, particularly histone methylation, which could drive the sustained transcriptional changes observed under transient hyperglycemia.

The role of chromatin modifications and epigenetic changes

The term ‘epigenetics’ was traditionally employed to describe heritable changes in gene expression and cellular phenotype attributable to mechanisms other than alteration of the underlying nucleotide sequence of the DNA [25]. Recently this definition has evolved to include the study of transcriptional regulation with a particular focus on chromatin architecture and the ensuing/preceding interactions between non-genetic factors and chromatin. Chromatin is a dynamic complex of DNA, histone proteins, and numerous modifying complexes, that serves not only to efficiently package DNA, but also to provide a mechanism of transcriptional regulation [25]. Eukaryotic gene transcription is a precisely regulated and multifaceted process, and among other mechanisms is primarily controlled by
the degree of chromatin accessibility. Altered states of transcription are initiated or perpetuated by the dynamic structural adaptation of regions of chromatin to confer transcriptionally permissive or repressive configurations [26]. This in turn regulates access and subsequent enzymatic activity of the core transcriptional machinery and associated factors to the underlying DNA sequence. Consequently, the structural state of chromatin determines the transcriptional competency of a gene at a particular locus [27]. Environmental factors are proposed to significantly influence the epigenetic signature, and therefore the transcriptional competency of chromatin. Emerging evidence suggests a key role for deregulated epigenetic transcriptional control in the pathogenesis of numerous human diseases including metabolic and inflammatory disorders associated with diabetes. The cellular transcriptional response to various environmental is exemplified by extensive reports of epigenetic transcriptional changes in response to various stimuli and insults in cell culture and experimental models. Thus exposure to environmental factors such as hyperglycemia may lead to the establishment of altered transcriptional states of chromatin that could potentially be retained for the lifetime of the organism. Furthermore there is increasing evidence to indicate that in simple eukaryotes as well as animal models, epigenetic transcriptional states can be meiotically inherited and trans-generationally persistent [28,29].

**DNA methylation and diabetic complications**

A classical example of epigenetic transcriptional control is the covalent post-replicative modification of DNA by the addition of methyl groups to cytosine residues primarily at CpG dinucleotides within the genomic sequence, leading to transcriptional repression. Cytosine methylation within DNA regulatory regions can inhibit transcription by physically precluding the association of DNA and transcription factors [30] and influencing nucleosomal positioning [31,32]. Methylated CpG residues are recognised and bound by several methyl-CpG binding domain proteins which associate with various chromatin modifiers to establish transcriptionally repressed chromatin [33-35]. Limited studies have focused on the role of DNA methylation in the pathogenesis of diabetes, however altered methylation patterns have been reported in patients with diabetes [36].

Regions of mammalian promoters enriched for CpG dinucleotides (CpG islands) are subject to dynamic methyl modification during development [37]. Studies suggest that *in utero* exposure to a poor nutritional environment might predispose the fetus to the development of diabetes in adult life through aberrant DNA methylation in pancreatic islets of rats [38] and at
the HNF4A gene locus of CD34+ stem cells from cord blood of neonates [39]. Importantly
DNA methylation has been shown to play a role in regulation of insulin expression. Analysis of
mouse embryonic stem cells revealed that DNA at the insulin (INS) promoter was methylated,
and became demethylated as the cells differentiated into insulin expressing β-cells. Observation of this specific demethylation extended to the human insulin promoter of pancreatic β-cells [40].

Several studies have examined the potential role of DNA methylation in the development of
diabetic complications, however the significance remains to be completely understood. One
study demonstrated global DNA hypermethylation of peripheral blood leukocytes from
patients with chronic kidney disease [41]. However analysis of kidney cells exposed to
hyperglycemia and renal tissues from Type I diabetic rats showed no difference in DNA
methylation of several candidate gene promoters [42]. Conversely numerous candidate genes
linked to atherosclerosis were revealed to have altered DNA methylation patterns in
endothelial cells [43]. With increased availability of high-throughput technologies for
examining the DNA methylome, several observations relevant to diabetic complications have
been highlighted by genome-wide studies. Genome-wide analysis of peripheral whole blood
DNA methylation patterns from a cohort of 192 Type I diabetics recently identified 19
prospective CpG sites associated with the risk of diabetic nephropathy [44]. Global analysis of
DNA methylation patterns in peripheral blood monocytes revealed significant changes in
patients with increased risk for cardiovascular disease [45]. Furthermore, studies in smooth
muscle cells of human atherosclerotic lesions and animal models including ApoE-null mice fed
high-fat diets revealed associations between atherosclerosis and global DNA hypomethylation
[46-48]. With regard to the development of Type I diabetes, recent genome-wide analysis of
DNA methylation of monocytes from monozygotic twins discordant for Type I diabetes
identified several diabetes-specific methylation variable positions. Further analysis revealed
that the methylation patterns of some of these positions were altered in individuals prior to
diagnosis. This suggests that they arise early in the pathology of Type I diabetes and unlike
aforementioned examples cannot be explained by post-disease cellular dysfunction [49].

Chemical diversity of histone tail modifications
Traditionally considered as primarily passive structural elements histone proteins are now
recognised as integral components of chromatin fundamental to the regulation of gene
expression [50,51]. Underlining their importance to nuclear regulation, histones are among
the most highly conserved proteins through the eukaryotic domain [52]. Each histone peptide possess an unstructured N-terminal tail composed of 15-44 amino acids that protrude from the nucleosome to provide exposed surfaces for interaction with other proteins. The tail domains harbor multiple lysine and arginine residues that are substrates for a variety of covalent post-translational modifications including methylation, acetylation, phosphorylation and ubiquitination [53]. Many of these modifications are associated with distinct transcriptional states and can generate synergistic or antagonistic effects in conjunction with other histone modifications to control transitions between transcriptionally active and inactive chromatin [27].

The wealth of post-translational modifications to the histone tail suggests a complicated and combinatory epigenetic language that co-ordinates structural and therefore transcriptional changes. Indeed several observations indicate that acetylation of particular histone tail residues can complement the effects of other histone modifications such as methylation on gene transcription [54]. Enzymatic transfer of an acetyl group to the ε-amino group of histone H3 and H4 lysine tails by histone acetyltransferases (HATs) has been correlated with transcriptional activation [55] most likely through alterations of electrostatic interactions [56]. By contrast, removal of the acetyl group by histone deacetylases (HDACs) is associated with transcriptional repression. Accordingly euchromatin is highly enriched for acetylated histones, whereas hypoacetylated histones are observed predominately at transcriptionally inactive heterochromatin [57,58]. Such broad observations have been proposed to reflect transcriptional competency rather than active transcription of such regions [59]. Nonetheless, hyperacetylation of certain histone lysine residues, particularly K9 and K14 of histone H3, has been thoroughly demonstrated as characteristic of promoter and enhancer regions of genes in active transcriptional states and several enzymes and complexes responsible for the addition and removal of the acetylation mark have been described [60-63]. Recent studies have explored the relationship between histone hyperacetylation and gene expression under diabetic conditions. Cultured primary human endothelial cells exposed to hyperglycemic conditions displayed increased expression of p300 acetyltransferase and enrichment of this enzyme to the promoters of HG-responsive genes such as fibronectin [64]. Hyperacetylation of histone H4 at the promoter enhances expression of the INS gene [65]. Furthermore the β-cell specific transcription factor PDX1 was shown to interact with p300 to mediate proinsulin gene expression [66]. Thus histone acetylation is involved in both regulation of insulin secretion and damage associated with hyperglycemia.
Therapeutic targeting of the enzymes that regulate these processes may therefore alleviate some aspects of the disease. To this end the anti-inflammatory effects of HDAC inhibition have recently been investigated. Genome-wide mRNA sequencing of human aortic endothelial cells revealed decreased expression of numerous inflammatory cytokines and chemokines such as IL-6, IL-8, MCP-1 and MIF and cell adhesion molecule ICAM-1 upon treatment with the HDAC inhibitor Trichostatin A (Balcerczyk et al., manuscript in preparation). The aim of this review is not to provide an exhaustive description of each histone modification, which have recently been reviewed [67], but instead we focus on key relevant histone modifications in context to hyperglycemia, metabolic memory and its persistence. Perhaps the best characterized enzyme relevant to diabetic complications is the Set7 methyltransferase in primary culture experiments and small animal models of hyperglycemic memory.

Histone methylation

Histone methylation modifications catalysed by histone methyltransferases (HMTase) define and direct the formation of distinct chromatin regions [68,69]. Contrasting histone acetylation, methylation of lysine and arginine residues can be associated with both gene activation and repression, depending on the residue modified [69]. Most HMTase enzymes that specifically methylate lysine residues are classified as members of the SET family of proteins. Responsible for the methyltransferase activity is the presence of the evolutionarily conserved, 130-amino acid SET domain initially identified in three proteins shown to be required for maintenance of the expression of genes important to normal development in Drosophila melanogaster; positive effect variegation suppressor SU(VAR)3-9, the polycomb group protein Enhancer of zeste and the homeobox gene regulator Trithorax [70-73]. Identification of this common motif has rapidly increased the number of genetically and biochemically identified modifiers of histone proteins across several organisms. Multiple SET-domain HMTases have been characterized that methylate histone lysine residues in mammalian cells (Table 1). SET domain HMTases contribute to the regulation of transcriptional activity by methylating histone proteins within chromatin to maintain active and repressed transcriptional states through various nuclear processes. These modifications occur on several lysine residues within the N-terminal tail of histones H3 and H4, and the effect on transcription is dependent on both the specific lysine residue modified and the number of methyl groups covalently assigned. The enzymes responsible generally exhibit strong substrate and product specificity. Methylation of lysine residues 4, 36 and 79 of histone H3 are associated with active gene expression [25,74,75], while methylated lysine
residues 9 and 27 of histone H3 and lysine 20 of histone H4 are generally associated with transcriptionally silenced or inactive chromatin [25,75,76]. Recent genetic examination of genes encoding histone lysine methyltransferases in patients with Type 1 diabetes suggests the importance of an inherited genetic component for the risk of diabetic complications. In the well-characterised Finnish Diabetic Nephropathy Study (FinnDiane) cohort, a polymorphism in the SUV39H2 gene that encodes a H3K9-specific methyltransferase was found to be associated with diabetic retinopathy [77]. Though this gene has not previously been linked to diabetic complications, it is a close homologue of SUV39H1 that has been implicated in hyperglycemia-induced inflammation [17].

<table>
<thead>
<tr>
<th>Histone Target</th>
<th>Lysine methyltransferase</th>
<th>Monomethylation</th>
<th>Dimethylation</th>
<th>Trimethylation</th>
</tr>
</thead>
<tbody>
<tr>
<td>H3K4</td>
<td>Set7</td>
<td>Smyd3, SETMAR, NSD3</td>
<td>Smyd3, MLL, SETD1A, SETD1B</td>
<td></td>
</tr>
<tr>
<td>H3K9</td>
<td>G9a, Eu-HMTase1</td>
<td>G9a, Eu-HMTase1</td>
<td>SETDB1, SETDB2, SUV39h1, SUV39h2</td>
<td></td>
</tr>
<tr>
<td>H3K27</td>
<td>EZH2, EZH1, G9a</td>
<td>EZH2, EZH1, G9a, NSD3</td>
<td>EZH2, NSD3</td>
<td></td>
</tr>
<tr>
<td>H3K36</td>
<td>NSD1, ASH1L</td>
<td>NSD1, Smyd2, SETMAR, ASH1L</td>
<td>SETD2, ASH1L, NSD2</td>
<td></td>
</tr>
<tr>
<td>H3K79</td>
<td>DOT1L</td>
<td>DOT1L</td>
<td>DOT1L</td>
<td></td>
</tr>
<tr>
<td>H4K20</td>
<td>Pr-Set7(Set8), NSD2</td>
<td>Pr-Set7(Set8), NSD1, SUV4-20</td>
<td>NSD2, SUV4-20</td>
<td></td>
</tr>
</tbody>
</table>
Set7 methyltransferase

A SET-domain HMTase has recently received considerable attention for its ability to catalyse the activating H3K4 monomethylation (H3K4me1) reaction. Sharing little sequence homology with any of the six previously described yeast SET-domain containing proteins, the enzyme was termed Set7 [78] (also described as Set9) [79]. Initial characterization demonstrated the ability of Set7 to stimulate GAL4-VP16-activated luciferase expression in cancer cell lines, implying a role in activator-induced transcription [79]. To this end, several functions attributable to Set7 activity have been observed in vitro which may be important to understanding the mechanism behind the transition between transcriptionally inactive and active chromatin. Set7-mediated H3K4 methylation inhibits Suv39h1-mediated H3K9 methylation and facilitates acetylation of both H3 and H4 by the acetyltransferase p300 [80]. Furthermore, association of the NuRD chromatin remodeling and deacetylase complex with the histone H3 tail is inhibited by H3K4 methylation, but not H3K9 methylation [79]. This antagonistic paradigm suggests that Set7-mediated H3K4me1 has the ability to maintain an active chromatin state by preventing both NuRD- and HP1-mediated (H3K9 methylation-dependent) transcriptional silencing. Interestingly, methylation of H3K4 by Set7 did not have the same effect on H3K9 methylation by G9a [79]. This observation excludes Set7-mediated H3K4me1 as a global regulator of H3K9 methylation and by extension global transcriptional repression.

Set7 methyltransferase regulates metabolic memory and inflammatory diabetic complications

Several studies associate Set7 methyltransferase activity at H3K4 with gene activation under diabetic conditions in pancreatic cells [81-83], monocytes [84] and endothelial cells [17,85]. Recently HG concentrations were demonstrated to promote nuclear localization of Set7 in human endothelial cells [86]. Reduced H3K4me1 enrichment at discrete hyperglycemia-responsive loci following Set7 depletion has been reported in models of hyperglycemic variability. Following transient exposure of endothelial cells to HG, persistent transcriptional activation of p65 (encoded by the RELA gene) and several inflammatory NFkB-B-dependent genes paralleling enrichment of Set7 and H3K4me1 at the RELA promoter was observed upon subsequent return to normoglycemia. This response was sustained for 6 days implying a memory of recent exposure to the metabolic insult. Set7 knockdown abrogated RELA promoter H3K4me1 enrichment and p65 mRNA and protein expression in response to...
transient and prior HG in both human and bovine endothelial cells. Accordingly the transcriptional persistence of p65 and its downstream transcriptional targets was attenuated [17,85]. Furthermore, sustained reduction of H3K9 methylation on the RELA promoter following transient and prior hyperglycemia was observed [85]. Methylation H3K9 and H3K4 have previously been demonstrated to inhibit each other [78] and the occlusion of H3K9 methylation by Set7-mediated methylation of H3K4 may represent another mechanism by which Set7 maintains an active chromatin state at the RELA promoter [85].

Experimental evidence suggests that Set7 may function as a co-activator of transcription through direct interaction with another transcriptional activator and specific co-recruitment to promoters. Monocyte transcription of a subset of TNF-α-inducible NFκB-dependent inflammatory genes was compromised by depletion of endogenous Set7 through a different mechanism than previously reported for glucose-stimulated endothelial cells. Expression of TNF-α-inducible p65-dependent MCP-1, IL-8 and TNF-α was shown to be dependent on Set7 expression [84]. TNF-α-stimulated cells displayed increased MCP-1 expression concomitant with enrichment of Set7, p65 and H3K4 methylation at the MCP-1 and TNF-α promoters. Targeted Set7 disruption using shRNA impaired this response, namely the promoter-specific recruitment of p65. Microarray analysis of TNF-α-stimulated Set7 knockdown monocytes revealed altered transcriptional induction of over 25% of TNF-α-regulated genes. Interestingly, the level of p65 expression detected by western blot was similar across control and Set7 deficient cells [84]. This observation indicates that the effects on downstream transcription targets were not a result of attenuated RELA transcriptional activation, as was the case in the response of endothelial cells to glucose stimulation. Rather Set7 appears to act as a co-activator of a subset of p65-mediated transcription of TNF-α-sensitive genes. While these seemingly discordant findings may reflect differences across cell types, it is likely that Set7 has a dual function in establishment of the pro-inflammatory phenotype.

Intriguingly the p65 subunit was recently identified as a novel substrate for Set7-mediated methylation. Studies have demonstrated this modification to have implications for p65 transactivity and downstream transcription of NFκB-dependent genes. This is a significant finding given the enzymes’s reported role in persistent transcriptional activation of the RELA promoter in endothelial cells, and further implicates Set7 in regulation of NFκB-dependent inflammatory pathways. A recent report describes complete abolition of in vitro p65 methylation by a single point mutation at K37, indicating this residue to be the lysine targeted
for modification [87]. Methylation dependent regulation of p65 may represent an integral control point with regard to the expression of a variety of genes.

While several known examples of posttranslational p65 modification exist, the biological response to Set7-mediated methylation of p65 remains to be fully established. Relative to the unmodified protein, p65 methylated at K37 is restricted to the nucleus and enriched at a subset of NFκB-dependent promoters in response to TNF-a stimulation [87]. Conversely, examples of this modification have also linked Set7 to transcriptional repression. Two alternative sites at K315 and K316 that are subject to Set7-dependent methylation have been reported, modification of which can result in negative regulation of p65 transactivity in vivo [88]. Subsequent kinetic analysis reveals that the K37 methylation event occurs prior to K315 and K316 methylation, and may be the preferential methylation site [87]. Nevertheless p65 appears to contain Set7-dependent regulatory sites for both transcriptional activation and repression of downstream gene targets. Overall Set7 appears to be a key mediator of the pro-inflammatory response to hyperglycemia through both increased p65 expression via promoter H3K4 methylation enrichment [17] and post-translational methylation of distinct lysine residues within the p65 subunit of NFκB [87,88]. In support of Set7-dependent regulation of p65 transactivity, recent massive parallel sequencing analysis of a model of Set7 depletion in human endothelial cells implicates deregulated p65 transactivity in the differential expression of numerous genes, some of which have associations with inflammation and diabetic complications (unpublished data) (Figure 1).

A similar co-activational role has been described in pancreatic β-cells with regard to Set7 recruitment and enrichment of H3K4 methylation at the INS promoter. Co-immunoprecipitation and GST pull-down assays in conjunction with ChIP assays demonstrate the recruitment of Set7 and subsequent methyltransferase activity to be mediated through direct interaction with transcriptional activator; pancreatic duodenal homeobox-1 (PDX-1). Depletion of either Set7 or PDX-1 expression significantly decreased transcription of a reporter construct driven by an intact INS promoter [83]. Significant reduction in transcription of a subset of glucose responsive genes was observed in mouse β-cells and primary islets in vitro, following siRNA-mediated Set7 depletion [81]. Unexpectedly, this transcriptional diminution correlated with reduced H3K4me2 enrichment at the proximal promoter regions of Ins1, Ins2 and Slc2a2, while no change to the H3K4me1 status was reported. The authors noted that these results do not necessarily rule out Set7’s function as a
monomethylase and raise the possibility of another closely linked HMTase capable of completing the di-methylation reaction on the enzymatic product of Set7 [81]. Importantly, the overall experimental observations highlight the necessity of Set7 function in β-cells for proper insulin production and secretion.

A recent study implicates Set7 as a co-activator of TGF-β-induced expression of extracellular matrix genes relevant to the development of diabetic nephropathy [89]. Under hyperglycemic conditions, increased expression of fibrotic genes Col1a1, CTGF and PAI-1 in rat mesangial cells was demonstrated to correlate with enrichment of promoter H3K4me1 and Set7 recruitment. Increased gene expression and Set7 recruitment were attenuated by both pre-treatment with a TGF-β-specific antibody and targeted Set7 disruption by siRNA transfection [89]. Col1a1 [90], CTGF [91] and PAI-1 [92] are bona fide transactivation targets of SMAD3, a key mediator of renal fibrosis and inflammation [91,93,94]. Indeed deletion of SMAD3 conferred protection against fibrogenesis in a rodent model of diabetic nephropathy [95]. Under diabetic conditions, critical mediators of diabetic complications AGEs [91,96] and Angiotensin II [93,97] can also activate SMAD signaling pathways to stimulate extracellular matrix production via TGF-β-dependent and independent mechanisms. Thus it is tempting to speculate a mediatory role for SMAD3 in the recruitment and specific enrichment of Set7 to activating genes targets in response to TGF-β stimulation to potentiate the H3K4me1 reaction. Interaction between Set7 and SMAD3 has not been demonstrated experimentally, but may provide another example of transcription factor-Set7 co-activation possibly via a methylation-dependent mechanism. Disruption of a potential Set7/SMAD3 interaction may provide a novel therapeutic target to alleviate renal complications associated with the diabetic milieu. Taken together, these findings provide evidence of co-recruitment of Set7 to specific genes and suggest that Set7 may function in complex with transcriptional co-activators. For instance ordered recruitment of Set7 and other transcriptional regulatory proteins indicate that Set7 co-assembles as a component of a pre-initiation complex required for transcriptional activation of collagenase I [98].

Set7 methylates non-histone substrates

As highlighted by reports of Set7-dependent post-translational methylation of p65 lysine residues, the molecular events mediated by Set7 in response to hyperglycemia might not be restricted to histone methylation. Analysis of Set7 methyltransferase activity toward lysine residues of non-histone proteins has revealed previously unknown mechanisms by which
Set7 may contribute to transcriptional regulation and other cellular processes [99]. Numerous additional substrates have recently been described sparking considerable interest in the characterization of these methylation events and the identification of novel lysine targets [100]. Adding to the complexity of the role of Set7 in transcriptional regulation, transcription factors predominate the list of confirmed Set7 substrates. The in vivo consequences of post-translational methylation on most of these targets remain to be fully understood. However numerous observations of Set7 methyltransferase activity-dependent transcription factor stability and activity have been reported, and several examples are described in Figure 2.

Recent studies suggest novel mechanisms of Set7 recruitment to individual promoters by Set7-transcription factor binding and modification, providing further evidence of a co-activational role. Several recently identified Set7 substrates display modified biological activity following Set7-mediated methylation including p53 [101], DNMT1 [102] and ER-α [103]. This post-translational methylation can result in altered expression of downstream transcriptional targets. For instance the TFIID transcription factor complex component TAF10 is methylated at K189 by Set7 [104]. While this modification does not affect the incorporation of TAF10 into TFIID, methylated TAF10 displays increased affinity for RNAPII suggesting a role for this modification in regulating preinitiation complex formation. To this end, transcription of several known TAF10-dependent genes was enhanced by methylation of TAF10 [104]. While this report highlights an example mechanism whereby Set7 is brought within close proximity of an activating promoter, subsequent effects on H3K4 methylation remain unresolved.

Sequential Set7 recruitment to histone H3 following TAF10 promoter binding remains to be demonstrated. Recent reports suggest this indeed to be the case with an interaction between Set7 and another recently characterized substrate. STAT3 is methylated at K140 by Set7, and de-methylated by LSD1 [105]. This modification both positively and negatively regulates the expression of a multitude of STAT3-dependent genes in response to IL-6 stimulation, and can be disrupted by Set7 depletion or cells expressing STAT3 R140 mutant proteins. Furthermore, STAT3 promoter occupancy of a subset of these genes was shown to be dependent on Set7 methyltransferase activity. This regulation is promoter specific as not all STAT3-dependent IL-6 responsive promoters exhibited methylated STAT3 enrichment [105]. IL-6 is considered to play a central role in the development of diabetic complications
including diabetic nephropathy [106,107]. Circulating IL-6 levels are reportedly elevated in patients with type 2 diabetes [108], are associated with poor glycemic control [109]. Recent microarray analysis of diabetic kidneys revealed significant up-regulation of genes in the JAK/STAT pathway within glomeruli and tubules [110]. Furthermore components of the diabetic milieu including ROS [111], angiotensin II signaling [105] and AGEs [112] have been experimentally demonstrated to activate this pathway in kidney cells. In a study of diabetic mice with compromised STAT3 activity, extracellular matrix-associated TGF-β and type IV collagen expression as well as mesangial cell proliferation were inhibited. Attenuation of the diseased phenotype extended to a reduction in the expression of pro-inflammatory factors IL-6, ICAM-1, MCP-1 and nuclear translocation of NFκB [113]. Given the responsivity of Set7 to hyperglycemia and ROS and its transactivity-modifying capacity of STAT3, it is reasonable to speculate a role for the methyltransferase in the development and progression of diabetic nephropathy through this pathway.

The SOCS3 gene was strongly up-regulated following Set7 knockdown, and displays increased STAT3 enrichment at the promoter [105]. Furthermore it was demonstrated that Set7 is recruited to the SOCS3 promoter after STAT3 recruitment. This sequence of events may function to clear STAT3 from the activating promoter and simultaneously potentiate methylation of H3K4 [105]. Set7/STAT3-regulated expression of SOCS3 may hold therapeutic relevance for diabetic nephropathy as SOCS3 is a negative-feedback regulator of the JAK/STAT pathway [114], specifically STAT3 activation [115]. To this end, over-expression of SOCS3 inhibited the hyperglycemia-induced activation of STAT-responsive inflammatory genes MCP-1, ICAM-1 and IL-6 [116].

A similar interaction between Set7 and the androgen receptor (AR) has been described [117] as a methylation event important for AR activation and recruitment to androgen-regulated promoters. This interaction may facilitate H3K4me1, as depletion of endogenous Set7 in cancer cells reduced H3K4me1 enrichment at proximal and enhancer regions of the androgen-dependent PSA promoter [117]. This observation holds relevance for diabetic complications, as hyperandrogenemia is associated with β-cell dysfunction [118,119]. In support, a recent study in a mouse model reported increased systemic oxidative stress stemming from testosterone treatment was AR-dependent. Furthermore female mouse pancreatic islets treated with streptozotocin displayed increased β-cell destruction [120].
Further investigation may reveal Set7 as a potential therapeutic target in the treatment of hyperandrogenemia and the accompanying insulin resistance.

**Conclusion**

Estimates indicate that the incidence of diabetes mellitus and associated complications is rapidly increasing. It is therefore imperative that strategies to combat progression of detrimental complications are established. Knowledge of the molecular etiology of diabetes mellitus is paramount to understanding the link between environmental factors such as nutrition and altered/persistent gene expression that drive the disease phenotype. Clinical and experimental studies have revealed the deleterious consequences of exposure to hyperglycemia, and clearly highlight the importance of strict glycemic control. The concept that a cell retains a metabolic memory of prior exposure to hyperglycemic events was a seminal hypothesis that arose from epidemiological studies of diabetic patients. This concept is further exemplified by experimental studies that demonstrate hyperglycemia-induced changes in gene expression that persist despite a return to physiological glucose concentrations.

The dynamic processes that govern epigenetic regulation of chromatin confer an additional layer of transcriptional control that links gene expression with environmental signalling. Experimental evidence demonstrates that hyperglycemia can induce epigenetic modifications to chromatin structure via various nuclear processes. Specific histone-modifying enzymes have been implicated, and have been shown to regulate the expression of inflammatory genes in cells of vascular origin. In these models of glycemic memory, the Set7 methyltransferase was shown to engage the histone H3 tail to establish H3K4me1 at the pro-inflammatory RELA promoter, a modification that persists despite normalisation of glucose conditions. Certainly other histone modifications and the responsible enzymes are expected to be involved in transcriptional changes induced by elevated glucose concentrations, and are likely to be explored in models of glycemic memory. Characterisation of these events will be greatly assisted by recent advances in powerful genome-wide technologies such as massive parallel sequencing and bioinformatics tools. For instance genome-wide analysis of primary vascular cells exposed to HG concentrations has revealed relationships between acetylated histone tails, DNA methylation and expression of genes associated with endothelial dysfunction [121].
The role of Set7 in gene activational events under diabetic conditions extends to pancreatic and monocytic transcriptional regulation that also hold relevance for diabetic complications. Additionally, recent and rapid expansion of substrates alternative to the histone H3 tail may provide further evidence of the enzyme's involvement in the pathogenesis of diabetic complications. Defining non-histone Set7 substrates is critical to understanding the role of Set7 in transcriptional regulation in health and disease. This avenue of investigation may uncover novel mechanisms by which Set7 is recruited to activating promoters at specific loci to execute the H3K4 methylation reaction. By the same token, Set7-regulated protein activity and stability, and consequential effects on transactivation potential is another mechanism of transcriptional control that may be applicable to diabetic complications. To attenuate the burden of diabetic complications it may soon be possible to interrupt the epigenetic pathways that promote diabetic complications such as vascular injury. Specific targeting of the enzymes implicated in chromatin structure and gene function such as Set7 provides potential for future epigenetic therapy for diabetic complications. However several challenges to this approach are apparent in the multiplicity of modifications that govern gene expression. The combinatorial nature of histone modification patterns at individual loci and genome-wide will be rapidly unveiled with advances in sequencing technology. It is anticipated that this will strongly build upon key findings from *in vitro* studies of glucose- and ROS-induced vascular injury and potentially lead to therapeutic inhibition of methyl-writing and methyl-erasing enzymes to attenuate complications of diabetes.

Another strategy to abrogate the effects of hyperglycemic exposure is inhibition of ROS overproduction. Oxidative stress plays a pivotal role in the development of complications of diabetes by activating numerous pathogenic pathways [122]. Furthermore increased ROS production inactivates 2 athero-protective enzymes eNOS [123,124] and prostacyclin synthase [125]. The significant reduction of eNOS and prostacyclin synthase activity attributable to hyperglycemia-induced ROS overproduction was attenuated by treatment with a superoxide dismutase/catalase mimetic [23]. Similar strategies of ROS inhibition have been demonstrated to prevent the development of complications including cardiomyopathy [126,127], nephropathy [128,129], retinopathy [130] and neuropathy [131] in diabetic mice.
Figure 1 Massive parallel sequencing analysis of a model of Set7 depletion in human endothelial cells. Deregulated p65 transactivity is implicated in differential expression of numerous genes, some of which have associations with inflammation and diabetic complications. Vertical lines categorise the cellular location of protein products of the deregulated genes in the network: extracellular, membrane, cytoplasm, nucleus and unspecified. The single arrow going toward p65 marks the interaction with the Set7 methyltransferase. Red circles in the right corner of a gene indicate increased expression and blue circles indicate decreased expression. This transcription factor centric network was generated using the commercial software MetaCore™ version 6.8 build 30387 (GeneGo, Inc) which draws from the manually curated MetaCore database. 
Figure 2 Examples of transcription-associated protein substrates for the methyltransferase reaction catalysed by Set7. Several of these modifications may contribute to a positive transcriptional outcome. Set7-mediated methylation of K142 destabilizes DNMT1 protein by facilitating polyubiquitination and subsequent proteasome-mediated degradation [102]. By contrast methylation of p53 at K372 and subsequent acetylation by Tip60 acetyltransferase results in increased stability of the p53 protein, thereby influencing p53-mediated gene expression [132]. Methylation of p65 at K37 methylated p65 is restricted to the nucleus and is enriched at a subset of NFκB-dependent promoters in response to TNF-a stimulation. Methylation at K140 and K630 of STAT3 and AR respectively results in co-recruitment of Set7 to target gene promoters as a co-activator of transcription. STAT3 is reportedly di-methylated by Set7 [105].
**Sources of Funding**

The authors acknowledge grant and fellowship support from the Juvenile Diabetes Research Foundation International (JDRF), the Diabetes Australia Research Trust (DART), the National Health and Medical Research Council (NHMRC) and the National Heart Foundation of Australia (NHF). STK is supported by an Australian Postgraduate Award (APA). AE-O is a Senior Research Fellow supported by the NHMRC. Supported in part by the Victorian Government's Operational Infrastructure Support Program.

**Declarations**

None
References


129. **DeRubertis FR, Craven PA, Melhem MF.** Acceleration of diabetic renal injury in the superoxide dismutase knockout mouse: effects of tempol. Metabolism. 2007 Sep;56(9):1256-64.


