Epigenetic Cues Emerging from Lysine Acetylation
Chromatin-Mediated Transcription to Mitochondrial Functions

Abstract
The landscape of lysine acetylation has expanded from nucleus to a cellular-wide global event. The substrates of histone acetyltransferase (HAT) in addition to chromatin and transcription factors now include metabolic enzymes, cytoskeletal proteins, molecular chaperones, ribosomal proteins and nuclear import factors. Together, lysine acetylation is involved in regulating protein stability, chromatin-mediated transcriptional activation, subcellular localization and activities of metabolic enzymes. Growing number of investigations reveal that a significant number (~2000) of proteins are undergoing acetylation on either one or more than one lysine residues. Notably, our review underscores the dynamic nature of acetylated lysine site(s), which can become a recruitment site(s) for bromodomain-containing proteins or histone deacetyltransferases that ultimately facilitates cellular response to diverse internal and external environmental changes. Finally, our review emphasizes on the strength of small chemical molecules, which do not perturb the expression of endogenous proteins to address complexities of epigenetic modifications. These small molecules can serve as tools to enhance our mechanistic insights as well as have therapeutic potential to treat many diseases.

Keywords: Epigenetics; Lysine acetylation; deacetylation; Chromatin; Transcription; Metabolism

Abbreviations: HAT: Histone Acetyltransferase; HDACs: Histone Deacetylase; FDA: Food and Drug Administration

Introduction
Amongst 20 amino acids, lysine is the most versatile owing to its steric flexibility, which facilitate several chemical modifications, such as acetylation, butyrylation, crotonylation, formylation, hydroxylation, malonylation, methylolation, propionylation, ubiquitination, succinylation and sumoylation [1-14]. Though serine/threonine phosphorylation and lysine acetylation were discovered between 1959 and 1963 respectively, most investigations focused on understanding the functions of kinases during that time [15,16]. Two major studies that propelled unraveling the significance of acetylation include the discovery of coenzyme A and acetyl-CoA, followed by the evidence that acetylation of histone proteins could modulate gene transcription [17,18]. Growing number of studies have established that lysine acetylation plays a crucial role in the pathogenesis of diseases, including diabetes, obesity, inflammation and cancers [19-22]. Essentially, acetylation not only imparts gene regulatory capabilities to the transcriptional machinery, but it also governs cellular processes not limited to DNA replication, DNA damage and repair, chromatin remodeling, metabolism and cell cycle progression [23-27]. Hereafter, the notion that acetylation is restricted to modulating chromatin-mediated transcription function expanded and now, it is recognized to be a global event involving mitochondrial and cytosolic proteins [28].

Consequently, lysine acetylation is not only limited to delivering plasticity to the chemical landscape of chromatin, but it also affects stability, sub-cellular localization and molecular interactions of proteins [29]. Additionally, the physiological and cellular roles of auto acetylation remain to be fully understood [30]. Transition from investigating the role of acetylation during chromatin-mediated transcriptional regulation to deciphering the significance of acetylation on the functions of transcription factors, such as p53, established the epigenetic basis of gene regulation during cellular response to stress. The major impact of these findings navigated the elucidation of acetylation-mediated molecular interactions with bromodomain and dynamic regulation of acetylation/deacetylation by histone deacetylase (HDACs) [31] (Figure 1). Furthermore, the identification of NAD-dependent sirtuins in mitochondria indicated that acetylation could regulate metabolic events also [31]. In 2009, Mann and his colleagues using high-resolution mass spectrometry discovered that there are at least 3,600 acetylated lysine sites (Kac) sites in 1,750 cellular proteins suggesting acetylation of multiple lysine residues on a single protein [1,28]. Notably, about 500 acetylated proteins participate in the chromatin-templated processes. Out of which, approximately 50 acetylated proteins are most likely involved in the process of DNA replication and another 80 could potentially play a crucial role during the process of DNA damage and repair [1,28]. Similarly, about 40 acetylated proteins could be involved in regulating of cell cycle genes and approximately, 60 proteins in nucleotide exchange factors [1,28].

Depending upon cellular environment conditions, about 55 proteins, which are involved in DNA/RNA helicase activity and nearly, 56 proteins possessing ubiquitin ligase and deubiquinases activities could undergo acetylation [1,28]. Given that multiple lysine residues are acetylated on a protein raises the possibility of a combinatorial pattern of acetylation, which could serve as a signature for facilitating cellular response to a diverse varieties of stress [28].
During gene transcription, one of the major challenges is to define acetylation-specific outcomes mainly because most histone acetyltransferases (HATS) are comprised of multi-domain proteins, which are capable of executing more than one function at a given time [32]. For instance, p300 and CBP are comprised of multiple cysteine-rich moieties, plant like homeodomain and bromodomain in addition to HAT. Similarly, MYST1 is composed of chromodomain that binds to methylated-lysine as well as a HAT domain which acetylate human histone H4 on lysine 16 [33]. TAFII250 coactivator contains double bromodomain, a HAT as well as Kinase domains [34]. Clearly, due to these multitude of activities associated with a particular HAT, overexpression as well as genetic knockout strategies may not fully convey the acetylation-specific outcomes. Furthermore, depending upon the alterations of external and internal environmental conditions, lysine acetylation could be mutually exclusive to methylation or ubiquitination. Collectively, these data underline that epigenetic modifications could be best studied by using small molecules, which does not perturb the expression of endogenous proteins. In this direction, the development of acetylation-specific antibodies has been one of the most effective tools that enhanced our understanding of the kinetics driving acetylation [35]. With the advent of proteomic and genomic high throughput technologies knowledge of new molecular targets that modulate gene functions have tremendously increased. Particularly, the dynamism exhibited by epigenetic modifications, such as acetylation/deacetylation, which if perturbed by a small molecule has potential to impact cell fate decisions that eventually has made them attractive targets for drug discovery program.

The FDA-approved HDAC inhibitor, Vorinostat, induces the differentiation of tumor cells has been approved for treatment of T cell lymphoma [36]. Besides, several inhibitors of bromodomain showed promising results in blocking pathogenesis of ischemia and HIV infection [37,38]. Most recently, inhibitor of BRD4, IPI140, which was tested in various model systems, suppresses cMYC by binding to the first bromodomain of BRD4 [39]. Clearly, emerging studies show promising small molecules that target HDACs or bromodomain, but a selective ligand for histone acetyltransferase remains to be developed. Most importantly, chemical biology is a gateway for not only dissecting the function of an endogenous protein with multiple functions but also could offer highly selective ligand which could have therapeutic value.

Taken together, the ability of amino acid lysine to facilitate quick chemical alterations that aid cellular capability to adjust as well as respond to environmental changes upon genomic stress, viral infection and energy generating metabolic reactions is remarkable. Given the genome-wide numerical ratio of HATs to HDACs or to Sirts as well as to bromodomains, it appears that we have only begun to scratch at the surface to understand the complete biological significance of acetylation in maintaining the cellular homoeostasis. However, we take this opportunity to salute investigators contributing through 50 years since the discovery of acetylation.

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