Bacterial nucleomodulins and cancer: An unresolved enigma

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A B S T R A C T

Recent studies in microbial pathogenesis have identified several bacterial proteins with the potential to influence host cell nuclei. This field of research is in its infancy, however it is rapidly growing. In particular, the role of bacterial nucleomodulins in animal oncogenesis is an area that requires attention. Earlier research has suggested the role of nucleomodulins in plant tumor development and these findings may provide us with a better understanding of the role of these proteins in human cancer development. This proposition is further supported by previous identification of nucleomodulins present in bacteria that have been associated with cancer development, but their role in human cancer is unclear. In this article, we provide an update on the status of these nucleomodulins and their role in cancer etiology. We collected information about known bacterial nucleomodulins and tried to relate their mechanistic implication with already known plant tumor development model. The present research indicates that bacterial nucleomodulins may be an important target in cancer etiology and knowledge of their role in human oncogenesis may help us to create suitable alternative cancer management strategies.

Introduction

While pathogens may be small, they have the superb ability to influence comparatively more complex, larger host cells. Evolutionary advantage made the pathogens advanced and efficient with multiple strategies to target host cells. They can secrete certain effectors to hijack host cellular and subcellular machinery for their survival. Bacteria are also known to release proteins that target important subcellular organelles including the nucleus, mitochondria, endoplasmic reticulum, Golgi body etc. Several reviews concerning the regulation of host cellular machinery by pathogen’s proteins through subcellular targeting are available and the list of such proteins is increasing rapidly [23,51]. Among these proteins, some have the ability to target and modulate nuclear function specifically. These proteins are known as nucleomodulins and have been shown to regulate nuclear function from inside to outside nucleus [9].

The nucleus serves as the core of the cell machinery and controls several important biological processes. It harbors genetic information and regulates expression of genes. The modulation of the nucleus by bacterial regulators, such as nucleomodulins, is important for controlling host cell machinery for bacterial advantage. Although, a deeper understanding of these nucleomodulins is needed and the detection of them in different bacteria is ongoing, several articles on the role of nucleomodulins in mediating host cell regulation arose recently [10]. The majority of literature surrounding bacterial nucleomodulins focuses on their role in host cell regulation and consequent contribution in disease pathogenesis, rather than their link to other mechanisms such as cancer development.

In fact, several bacterial infections are linked with cancer. It has been estimated that around 20% of cancer are attributed to infections [98]. There is substantial mechanistic involvement of microbial infections in the development of cancer, but exact mechanisms behind these findings are still disputed and studies investigating the mechanistic links between infection and cancer development are still ongoing.

Bacterial nucleomodulins may be an important player in cancer development due to their ability to alter nuclear function. Many reviews detail the role of bacterial nucleomodulins in host cell modulation, but their subsequent contributions in cancer etiology remains unexplored. Therefore, this article attempts to provide a glimpse into current status of bacterial nucleomodulins and their subsequent effects on cancer development through microbial involvement. Strong evidence of plant tumor development by bacterial nucleomodulins are currently available, but the similar role in human cancer development remains unclear.

Bacterial nucleomodulins and their consequences on host cell

As the nucleus plays a central role in cellular functions, modulation of nuclear function will ultimately lead to substantial effects on the host cell. Nucleomodulins are considered to be bacterial factors that target the nucleus [9,10]. Several articles are already available on the regulation of host cell by nucleomodulins so we briefly covered this aspect in
| Sr. No. | Name of Protein | Organism | Function (TSS) |
|---------|----------------|----------|---------------|
| 1       | VirE2 [97]     | A. tumefaciens | It is involved in host nuclear uptake of bacterial ssDNA element known as transferred DNA and results in DNA transformation (T4SS). |
| 2       | VirE3 [27]     | A. tumefaciens | Plant transcriptional activator (T4SS). |
| 3       | VirD5 [86]     | A. tumefaciens | VirD5 is known to have a dual function. It acts as a transcriptional activator and regulates host gene expression, it also prevents the degradation of coat proteins of the bacterial T-complex through the host’s ubiquitin proteasome system [86] (T4SS). |
| 4       | T-DNA border endonuclease virD2 [60] | A. tumefaciens | It is involved in the cleavage of unique sites within T-DNA of bacteria and involves the transfer and integration of particular segments (T-DNA) of the Ti plasmid DNA into host. T4SS effector (T4SS). |
| 5       | Protein VirF [80] | A. tumefaciens | In the host cell, VirF is involved in ubiquitination and subsequent proteasomal degradation of target proteins. It is also involved in T-DNA translocation and is essential for tumor formation in some plant species (T4SS). |
| 6       | Transcription Activator-Like (TAL) effectors [38] | Xanthomonas oryzae pv. oryzae | Functions as a transcription factor in rice plant cells (T3SS). |
| 7       | XopD [36]      | X. campestris | Interferes with host proteins regulation by mimicking host SUMO isopeptidase (T3SS). |
| 8       | AvrXa7 [89]    | Xanthomonas oryzae pv. oryzae | Bind to host double-stranded DNA and affects transcriptional machinery (T3SS). |
| 9       | HsvC [14]      | Pantoea agglomerans pv. gyropophilae (Pag) | DNA binding protein that controls expression of host gene HSVGT (T3SS). |
| 10      | HsvB [14]      | Pantoea agglomerans pv. gyropophilae (Pag) | Act as a transcriptional activator (T3SS). |
| 11      | HopA11 [14]    | Pseudomonas syringae | Proposed to Affect host gene expression by modulating post-translational modification of histones (T3SS). |
| 12      | PopP2 [14]     | Ralstonia solanacearum | Modulates host cell gene transcription (T3SS). |
| 13      | SET Domain protein (The Chlamydia nuclear effector [NUE]) [65] | Chlamydia trachomatis | Leads to host cell chromatin modification (T3SS). |
| 14      | hypothetical protein CT311 [53] | Chlamydia trachomatis | ?? (sec-dependent pathway) |
| 15      | Nuclear Effector AnKA [26] | Anaplasma phagocytophilum | Binds to host AT rich DNA and affects transcription of antimicrobial defense genes in granulocytes [71] (T4SS). |
| 16      | p200 [95]      | Ehrlichia spp. | Binds to adenine-rich DNA and affects host cell gene transcription (T1SS). |
| 17      | TRP12 [25]     | E. chaffeensis | Binds to G-rich motif of host DNA and modulates gene transcription (T1SS). |
| 18      | TRP47 [46]     | E. chaffeensis | Enters host nucleus through MYND binding domain and regulates transcription of essential genes (T1SS). |
| 19      | TRP120 [94]    | E. chaffeensis | Binds to GC-rich motif and shows transcriptional activator activity (T1SS). |
| 20      | OsPF [4]       | Shigella flexneri | Affects histone protein phosphorylation and host immunity-related genes expression (T3SS). |
| 21      | OsPB [99]      | S. flexneri | Modulates MAP Kinase pathway and affects host inflammatory response (T3SS). |
| 22      | E3 ubiquitin-protein ligase ipaH9.8 [5] | S. flexneri | Affect Nf-kb-mediated inflammatory response (T3SS). |
| 23      | E3 ubiquitin-protein ligase SipH1 [31] | Salmonella enterica serovar typhimurium | Inhibits Nf-kb associated genes (T3SS). |
| 24      | Cycle inhibiting factor (Cif) [41] | E. coli | Induces accumulation of cyclin-dependent kinase inhibitors p21 and p27, and inhibits cell cycle at both G1/S, G2/M phases (T3SS). |
| 25      | EppF [21]      | Entero-pathogenic E. coli | Disrupts host cell nucleolus and many other cellular events (T3SS). |
| 26      | YpHm [7]       | Y. pestis | Interacts with p90 ribosomal S6 kinase 1 (RSK1) and helps in virulence (T3SS). |
| 27      | Uncharacterized protein (CifYp) [19] | Y. pseudotuberculosis | Cell Cycle inhibiting factors (T3SS). |
| 28      | CufBp [68]     | Burkholderia pseudomallei | Induces methylation of host cell chromatin H3K4 protein and activates ribosomal DNA transcription (T3SS). |
| 29      | BiSET [23]     | Burkholderia thailandensis | Induces methylation of histone H3K4 protein (T3SS). |
| 30      | CufP [10]      | Photobabus luminescens | Cell cycle inhibiting factors (T3SS). |
| 31      | BuSET [23]     | B. anthracis | Targets host cell histone protein H1 |
| 32      | RomA [23]      | Legionella pneumophila | Leads to methylation of histone H3 Lys 14 (T4SS). |
| 33      | AnkH [84]      | L. pneumophila | Interferes with host cell transcription (T4SS). |
| 34      | AnkX [91]      | L. pneumophila | Interacts with host nuclear protein PLEKHM1, manipulating inflammation (T4SS). |
| 35      | SmPL [74]      | L. pneumophila | Targets host RNA polymerase II and regulates host gene expression (T4SS) |
| 36      | LntA [50]      | Listeria monocytogenes | Targets host cell chromatin repressor BAHD1 and stimulates interferon genes. |
| 37      | SuATI [77]     | Theliteria annulata (Protozoa) | Function as a DNA binding protein with the ability to alter host cell phenotype. |
| 38      | Rv2966c [76]   | Mycobacterium tuberculosis | Interacts with host chromatin and affects gene expression. |
| 39      | Rv1988c [90]   | Mycobacterium tuberculosis | Interacts with host chromatin and affects histone methylation. |
| 40      | serine/threonine phosphatase (SP-STP) [1] | Streptococcus pyogenes | Induces apoptosis in infected cells. |

In order to provide necessary background. Our main intention is to discuss the possible role of bacterial nucleomodulins in human oncogenesis and a representative group of nucleomodulins are presented in Table 1 Nucleomodulins can act in a variety of ways in order to alter host cellular functions and some of these are presented below.

### Bacterial proteins acting as transcription factors

A number of articles are already available on the regulation of host cells by nucleomodulins through various mechanisms. Several bacteria are known to regulate host cell transcription through the secretion of a...
variety of proteins that either directly or indirectly affect nuclear functions. Some bacterial nucleomodulins can even act as transcription activators. For example, the VirE3 protein of Agrobacterium can bind to the plants cell-specific transcription factor, pBRp, which belongs to the TF2B family. VirE3 is also able to induce transcription in yeast after binding to DNA [27]. The same category of proteins was found in another plant pathogen Pantoaea agglomerans. It is known to secrete HsG and HsVB, two proteins that have two nuclear localization signal to target host nuclei and are known to induce transcription of certain genes in host cells [87]. In addition, there is a specific category of bacterial nucleomodulins known as Transcription Activator-Like (TAL) effectors, function as transcription factors for host cells and induce expression of certain genes (Table 1) [38].

**Histone modification and chromatin regulation**

Nucleomodulins can act as transcriptional regulators granting them with exclusive unhibited access to host genetic material. Eukaryotic genetic material is larger than prokaryotes and it is usually packed in a smaller, more limited nuclear space. The packing of DNA is done in order to ensure no disturbance to the vital properties of genetic material-like replication, transcription, repair, and chromosomal segregation. Histone and chromatin remodeling proteins maintain this tight packaging of nuclear DNA, known as chromatin. Several microorganisms, including bacteria, have the ability to regulate chromatin in their host cells in order to regulate transcriptional control of their desired genes. The modification of the chromatin structure by bacteria has been reviewed already and many bacterial nucleomodulins are known to target this process [29]. Numerous bacteria are known to secrete special SET domain containing proteins, that possess the ability to modify chromatin structure [2]. Table 1 also lists some bacterial nucleomodulins known to alter chromatin structure.

**Effects on cell cycle and DNA integrity**

In addition to the role of bacteria in chromatin regulation and subsequent control on transcriptional machinery, several bacterial nucleomodulins are able to damage DNA, affect DNA repair, as well as alter the cell cycle. Some bacteria are also known to secrete DNA damaging bacterial effectors. For example, E. coli is known to secrete a bacterial toxin colibactin, with the ability to cause DNA damage [20]. Another bacterial protein, cytolethal distending toxin (CDT) is known to have DNA damaging properties that induce cell cycle arrest. Many Gram negative bacteria are known to produce CDT. The role of bacterial proteins in host DNA damage has been further reviewed in detail in other articles [28].

**Nucleus invaders bacteria**

Some bacteria themselves are known to directly invade host cell nuclei and influence their morphology and functions. These bacteria are often protected from host cytoplasmic defense mechanisms due to this unusual target location. Intracellular bacteria are largely found to be associated with Protists, but their association with animals has also been observed. Holospora spp. are one of the more well-known intranuclear bacteria, and infect Paramecia. However, this infection does not always cause problems for Paramecia, instead it provides a selective advantage as Holospora induces expression of heart shock proteins, which results in increased cell survival. Much remains uncertain about intranuclear bacteria, but the current status has been reviewed in a recent article [75]. Intranuclear bacteria are also found to be associated with mammalian cells. For example Rickettsia rickettsia has occasionally been observed to undergo intranuclear growth in host cells [13]. Moreover, another bacterium associated with Rickettsia is found within the nuclei of human liver cell and is known as Orientia tsutsugamushi [66]. While it has been suggested that this unusual intranuclear colonization may be accidental [75], this bacteria is known to cause various nuclear injuries, degeneration, and in some instances clumping of nucleoproteins. This suggests their presence in the nucleus may not be completely unintentional since there are direct effects observed on nuclear morphology and its functions [66]. Another bacterium known as CC99 is known for its ability to infect Amoeba and is often known as Amoeba resistant bacteria. CC99 is thought to infect nuclei of human HeLa and U937 macrophage like cells [24]. The knowledge about intranuclear bacteria is still in its infancy and has not yet been identified properly. Future research directed towards such bacteria will shed more light on the role of these bacteria in their contribution to disease pathogenesis.

**Potential of same category proteins in oncogenesis**

Primarily cancer involves the dysregulation of cellular homeostasis caused by altered cell growth control. This altered cell growth is associated with several peculiar hallmarks, including constant proliferative stimuli, evasion of growth suppression mechanisms and cell death resistance, increased immortality, angiogenesis, invasion, and metastasis activity. It is considered that these hallmarks of cancer are as a result of genomic instability and inflammation [30]. The involvement of bacteria in genomic instability and inflammation is widely discussed but their exact role in cancer etiology remains under investigation. Bacterial nucleomodulins are thought to have significant potential to alter genetic regulation since bacteria themselves are a great contributor to inflammatory mechanisms inside host cells.

It is firmly believed that targets of nucleomodulins in host cells are directly involved in cancer etiology, and many studies are ongoing to utilize these targets and associated pathways for cancer management (Fig. 1). The role of bacterial nucleomodulins in cancer etiology is yet to be established, but the cancer causing ability of the targets of nucleomodulins provides several caveats. Regulation of transcription factors is associated with many diseases including cancer and the role of transcription factors in cancer etiology has been substantially studied. For example many cancer cells depend on the transcription factor c-Myc for growth, proliferation, cancer severity and their derived clinical effects. The role of transcription factors in cancer etiology has been reviewed numerous times [52]. Recently, the targeting of transcription factors in cancer management has also been reviewed [8,57,70].

The role of histone modification in cancer etiology has also been reviewed in many recent articles [6,73]. Post-translational modifications in histone proteins including methylation, phosphorylation and acetylation are found to be linked with cancer etiology. This modulation in epigenetic mechanisms controlling gene expression can lead to the development of cancer [85].

The effects of cell cycle modulation in oncogenesis are also known. Briefly, cancer is a disease involving uncontrolled cell division. The cell cycle is a multi-step process and each step is controlled by certain checkpoints. The well-known protein p53, is mainly involved in blocking the progression of the cell cycle through the G1 step by activating production of protein p21 which inhibits cyclin dependent kinase. Generally p53 prevents cell cycle progression before DNA repair. Even in some instances, the p53 induces cell apoptosis, if the DNA damage is severe. Mutations in p53 genes are evident in many cancers and its effects have been reviewed extensively [58]. For example, Li-Fraumeni syndrome is associated with germline mutations in p53 gene, and leads to inherited susceptibility to multiple cancers and also linked with altered cell cycle control [54]. In addition, the OrfX protein produced by L. monocytogenes interacts with the RybP gene in the nucleus. This RybP gene has the ability to regulate p53 and its expression is decreased in human cancer tissues. Therefore this suggests the potential role of bacterial nucleomodulins in p53 regulation [17,67]. Alongside cell cycle control modulation and its role in carcinogenesis, repair mechanisms as a result of DNA damage are also altered during the development of cancer. Several anti-cancer treatment strategies target these repair pathways for cancer management [34].
Bacterial nucleomodulins have great potential to contribute to cancer etiology by affecting several cancer targets, nevertheless their conclusive role in mammalian oncogenesis is yet to be established. Scattered evidence infers the carcinogenic potential of bacterial nucleomodulins, but the overall mechanistic link leading to oncogenesis is much awaited. The following section will attempt to gather pieces of literature concerning the role of bacterial nucleomodulins in mammalian carcinogenesis.

**Bacterial nucleomodulins and cancer etiology**

*Agrobacterium tumefaciens: a model for the involvement of bacterial nucleomodulins in tumorigenesis*

It was understood over 100 years ago by Smith and Townsend that Gram-negative soil bacterium *Agrobacterium tumefaciens*, induces tumors in plants. Later it was found that a large Ti plasmid is required for *Agrobacterium* to induce tumors [11]. Now, it has been widely established that *Agrobacterium* have a complete set of machinery that induces tumors in host cells. This machinery includes several proteins including nucleomodulins. In addition to nucleomodulins, it has a specific sequence of DNA known as T-DNA, which gets transformed into the host cell. T-DNA transformation in host cells acts as a transfer of machinery for generation of bacterial nucleomodulins inside host nucleus. T-DNA induces the production of excessive amounts of growth hormones inside the host cell, including auxin and cytokinin, and leads to the induction of tumors by the uncontrolled proliferation of host cells [72].

*Agrobacterium* is known to produce two categories of proteins, one is known as Ti-plasmid encoded virulence gene products (vir), and the second is the chromosomally-encoded virulence proteins (chv). Several articles are available on *Agrobacterium* pathogenesis and consequent tumor formation in host plants. Fig. 2 gives an overview of some nucleomodulins involved in the process, in order to get an idea behind role of bacterial nucleomodulins in plant tumorigenesis.

**Nucleomodulins and cancer associated bacterial infection**

Nucleomodulins are known to have the potential to alter normal cellular functions, which can contribute to cancer development. The possible strategies of bacterial nucleomodulins in mediating tumor development are shown in Fig. 1. In addition, Table 2 identifies nucleomodulins found in bacteria that are associated with cancer and therefore with the potential role in tumorigenesis. As the list of nucleomodulins is still in its infancy, the complete picture of nucleomodulin mediated effects on tumorigenesis can only be inferred for now, Fig. 3 provides mechanisms for the contributions of bacterial nucleomodulins to cancer etiology based on current evidences. The next section is intended to cover a few case studies of bacterial nucleomodulins that originate from cancer associated bacteria.

**Helicobacter pylori**

*H. pylori* is a proven example of bacterial involvement in cancer etiology. Epidemiological evidence suggests the involvement of this bacterium in the etiology of gastric carcinoma specifically [82]. It is known to produce several proteins with the ability to alter nuclear function and some of these proteins are briefly discussed in Table 2. Among these proteins, some are directly translocated to the nucleus while some indirectly influence nuclear function. Recent studies found some *H. pylori* proteins directly localized in the host cell nucleus and contributing to gastric cancer development. For example, *H. pylori* proteins, HP0425 and HP0059 contain nuclear localization signal allowing them to translocate to the host cell nucleus. These proteins have DNase activity which is an important factor for tumor development [47,48]. In addition to these nucleomodulins, CagA and VacA are the most widely acclaimed proteins found in *H. pylori* known to induce a variety of effects on the host cell leading to its suggested carcinogenic potential. CagA is encoded by CagA pathogenicity island, which also encodes the T4SS secretion system similar to vir genes found in *Agrobacterium*. The CagA pathogenicity island is present in highly virulent strains of *H. pylori* and is assumed to have been acquired by horizontal gene transfer events [18]. CagA is also involved in a variety of events not restricted to its host nucleus, including tyrosine phosphorylation by Src family kinase and induction of changes in cell structure leading to a hummingbird phenotype in cell culture [33]. Furthermore, CagA is known to localize to the inner membrane of host cell surface, but it can activate the ERK signaling pathway which influences the expression of several genes in the host nucleus, inducing cell proliferation and gastric cancer. CagA is also involved in nuclear accumulation of β-catenin, induction of the transcription factor NF-kB, which contributes to cancer progression [62,78]. In addition, CagA also induces DNA damage by increasing the expression of spermine oxidase [16]. It has been found that CagA has ability to target nucleus and small fraction of CagA is also detected in host cell nuclei [81] making it an important component of bacterial nucleomodulins involved in cancer. Another widely acclaimed *H. pylori* protein VacA localizes to cell mem-
**Table 2**

Nucleomodulins from bacteria either known or suspiciously involved in induction of tumor.

| Sr. No | Name of nucleomodulins | Bacteria | Function related to carcinogenesis | Type of cancer |
|--------|-------------------------|----------|-------------------------------------|----------------|
| 1      | VirE3                   | *Agrobacterium* | Transcription induction [27]         | Plant Tumor    |
| 2      | VirF                    | *Agrobacterium tumefaciens* | T-DNA translocation for tumor formation | Plant Tumor    |
| 3      | HsvG                    | *Pantoea agglomerans* | DNA binding protein, transcriptional activator [87] | Plant Tumor    |
| 4      | SET Domain protein (NUE) | *Chlamydia trachomatis* | Chromatin modification [65] | Cervical cancer |
| 5      | cnpSET                  | *Chlamydia pneumoniae* | Chromatin modification [59] | Lung cancer |
| 6      | Colibactin              | *E. coli* | DNA damage [83] | Colorectal cancer |
| 7      | Cytolethal distending toxin (CDT) | *E. coli* | DNA damage, cell cycle arrest [39] | Colorectal cancer |
| 8      | Gf                      | *E. coli* | Migrates to host cell nucleus and affects NEDD8 [41] | Colorectal cancer |
| 9      | HP0425                  | *H. pylori* | Translocates to nucleus DNase I-like enzymatic activity [47] | Gastric cancer |
| 10     | HP0059                  | *H. pylori* | Translocates to nucleus DNase I-like enzymatic activity [48] | Gastric cancer |
| 11     | SspH1                   | *Salmonella enterica serovar Typhimurium* | Translocates to nucleus and inhibits NF-kB dependent genes [31] | Hepatobiliary cancer |
| 12     | Rv3423.1                | *Mycobacterium tuberculosis* | A 8 Kda proteins acetyllating histone H3 at the K9/K14 positions [40] | Lung cancer |
brane in order to form anion conducting channels. It causes the release of cytochrome C from mitochondria, leading to apoptosis. In contrast, some forms of this protein are strongly associated with gastric cancer and it is assumed that all *H. pylori* virulence factors act together to drive cancerous transformation. Further, VacA is involved in the translocation of β-catenin into nucleus resulting in cell proliferation [61], their direct targeting to host nucleus is not known. The role of CagA in the induction of gastric cancer has been reviewed extensively and therefore, in this article we only specify the aspects involved in modulation of nuclear function.

**Escherichia coli**

The role of *E. coli* in colorectal cancer has been suggested in a number of sources including epidemiological, experimental and computational studies. However, the exact reason behind the involvement of *E. coli* in colorectal cancer etiology is still awaited [42,44,45,79]. Several nucleomodulins have been detected in *E. coli* and noted for their potential involvement in CRC etiology. Some of these nucleomodulins are discussed in Table 2. *E. coli* is known to produce a toxin named Colibactin, which has the ability to cause DNA damage in host cells [83]. In addition, the contributions of other nucleomodulins like cytotoxic necrotizing factors (CNF) and cytolethal distending toxins (CDT) are also subject to current research. An *in silico* study predicted several *E. coli* proteins with the ability to localize in their host cell’s nucleus [42], but the exact role of this category of proteins in *E. coli* mediated colorectal cancer etiology is still unknown.

**Chlamydia**

*Chlamydia* is another bacterial genus which has been noted for its involvement in cancer. While, contradictory evidence exists, the role of *C. trachomatis*, *C. psittaci*, *C. pneumoniae* is suggested in cervical cancer [96], ocular adnexal lymphoma [15] and in lung cancer [92], respectively. The exact role of *Chlamydia* in the etiology of cancer is still a matter of debate, but the presence of nucleomodulins in these organisms provides support for their etiological potential. Some *Chlamydia* spp. are known to have SET domain proteins with the ability to alter chromatin structure. The role of SET domain proteins in cancer etiology has already reviewed recently [37], and Table 1 and 2 cover a few SET domain proteins identified in a few *Chlamydia* spp. with the ability to alter host chromatin. In addition, some *Chlamydia* spp. are known to have several other proteins targeting the host nucleus during pathogenesis. CT621 is a *C. trachomatis* protein, which is involved in translocation into the host nucleus and cytoplasm through the type 3 secretion system [35]. SINC is a T3SS protein from *C. psittaci*, which translocates to the inner nuclear membrane of host cell [56]. Nevertheless, there are many other Chlamydial proteins that are yet to be investigated for their potential to target host nuclei. Perhaps future research will unveil the reasoning behind the intriguing involvement of Chlamydia in cancer etiology and
these nucleomodulins will serve as a link to fill the gap between the role of bacteria in cancer induction.

Salmonella

*Salmonella* is another debated bacterium suspected to be involved in cancer etiology. Many studies have linked the infection of *Salmonella* with hepatobiliary cancer risk [55]. *Salmonella* is also known to have proteins with the ability to target and alter host cell nuclear function. It has been reported that *Salmonella* also produces cytolethal distending toxin (CDT), which translocate into the host nucleus in order to cause DNA damage and chromatin fragmentation [69]. SphH1 is another *Salmonella* effector which translocate into the host nucleus and suppress NF-kB dependent gene expression [31], however its exact role in *Salmonella* mediated cancer risk must be evaluated as the role of several other *Salmonella* effectors is known in modulating cancer related events. For example, SopE is a T3SS protein from *S. typhimurium*, which has the ability to activate Rho GTPase (CDC42 & RAC1) resulting in cytoskeletal rearrangement and ultimately leading to the entry of bacteria into non phagocytic cells. In addition, SopE is involved in the activation of MAP kinase resulting in the activation of signaling pathways leading to altered nuclear function [32]. Similarly, another protein SopE2, is thought to have same ability of modulating nuclear function, but it has also ability to activate NF-kB. Moreover, SopB is involved in protection of host cell from apoptosis.

Other bacteria involved in cancer etiology or cancer-associated infections

*Mycobacterium tuberculosis* infection is common among lung cancer patients. Some studies suggest that pulmonary tuberculosis is associated with an increased risk for lung cancer [88], while some reports indicate that the immunocompromised status of cancer patients makes them more susceptible to tuberculosis infection [3]. Under both circumstances, tuberculosis is strongly associated with lung cancer. The protein Rv3423.1 found in *M. tuberculosis* is isolated from chromatin of human macrophages infected with these bacteria. It was found that this protein has ability to acetylate histone H3 [40]. The involvement of *Streptococcus* in the etiology of colorectal cancer is also a controversial point, but several studies have found an association between *Streptococcus* and colorectal cancer. It has been found that cell wall proteins of *S. bovis* can induce oncogenic transformation in the colon [22]. It is found to have a histone like protein HpIA, although its role has been proposed in the adherence to colorectal cancer cells [12]. Nevertheless, their histone like structure leaves scope for their inclusion in the category of nucleomodulins. Future research may reveal various novel nucleomodulins like properties of HpIA or even in other proteins found in *S. bovis*, as this bacterium has been identified for its ability to modulate nuclear function and its carcinogenic potential without identifying exact bacterial proteins (Table 2).

Concluding remarks

Host pathogen interactions span a very complex set of mechanisms involving a variety of molecules including nucleomodulins. Nucleus is the core of a cell and plays a very important role during oncogenesis. Hijacking of cell organelles, such as the nucleus, by pathogens is a very important aspect for understanding the role of bacteria in cancer etiology. While, many methods are available to detect nuclear localization of bacterial proteins and their subsequent effects on nuclear function of a certain proteins, they are extremely arduous for analyzing whole bacterial proteomes for their potential to contain nucleomodulins. Some computational tools are also available to alleviate this difficulty by predicting the nuclear localization of certain protein. However, while this process may be fast and convenient, recent research indicate that these tools also have some limitations when predicting bacterial protein localization in the nucleus of human cell [43]. In addition, some proteins can modulate nuclear function without translocating into the host cell nucleus making the situation more complex. Perhaps future research will identify alternative fast track methods to detect entire bacterial proteomes for their nucleomodulins potential and their subsequent impact on cancer etiology. Moreover, among Cif family, *E. coli* effector is known to target host nucleus, but much is not known about Cif from other bacteria and they are mentioned in text due to their homology with *E. coli* Cif and possible similar effects. Similarly, colibactin is known to cause DNA damage and nucleomodulation, but its physical nuclear targeting needs investigation. We included here several bacterial proteins with the ability to modulate nuclear functions, but their direct nuclear targeting needs more investigations. Therefore, more studies are needed to understand mechanisms of nucleomodulins homologs and other nuclear function modulators.

Despite these limitations, various findings are beginning to be made on the modulation of nuclear function by bacteria. A similar category of molecules is also frequently detected in cancer-associated bacteria and provides several implications. Bacterial nucleomodulins may be the perfect candidate for the study of the bacterial role in cancer etiology since the role of similar nucleomodulins found in plants in the induction of plant tumors has already been proven. Increased understanding of this mechanism and its applicability to human tumor development may open up many ways to manage cancer by controlling certain risk factors. Future research in this direction is needed to reveal the association between bacterial nucleomodulins and animal oncogenesis which will ultimately aid in the study of cancer associated microbes and their etiologic potential.

Declaration of Competing Interest

Authors declare no potential conflicts of interest related to text of this manuscript.

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