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The Polyamine Spermidine Modulates the Production of the Bacterial Genotoxin Colibactin

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ABSTRACT Colibactin is a polyketide/nonribosomal peptide produced by Escherichia coli strains that harbor the pks island. This toxin induces DNA double-strand breaks and DNA interstrand cross-links in infected eukaryotic cells. Colibactin-producing strains are found associated with colorectal cancer biopsy specimens and promote intestinal tumor progression in various murine models. Polyamines are small polycationic molecules produced by both microorganisms and eukaryotic cells. Their levels are increased in malignancies, where they contribute to disease progression and metastasis. In this study, we demonstrated that the endogenous spermidine synthase SpeE is required for full genotoxic activity of colibactin-producing E. coli. Supplying spermidine in a ΔspeE pks/H11001 E. coli strain restored genotoxic activity. Spermidine is involved in the autotoxicity linked to colibactin and is required for direct damaging activity on DNA. The production of the colibactin prodrug motif is impaired in ΔspeE mutants. Therefore, we demonstrated that spermidine has a direct impact on colibactin synthesis.

IMPORTANCE Colibactin-producing Escherichia coli strains are associated with cancerous and precancerous colorectal tissues and are suspected of promoting colorectal carcinogenesis. In this study, we describe a new interplay between the synthesis of the genotoxin colibactin and the polyamine spermidine. Polyamines are highly abundant in cancer tissue and are associated with cell proliferation. The need for spermidine in genotoxic activity provides a new perspective on the role of these metabolites in the pathogenicity of colibactin-producing E. coli strains in colorectal cancer.

KEYWORDS Escherichia coli, biosynthesis, colorectal cancer, genotoxic colibactin, polyamines

The genotoxin colibactin is a secondary microbial metabolite synthetized by Escherichia coli and other enterobacteria. The genetic determinant of colibactin is a 54-kb gene cluster, the pks genomic island (1). This highly conserved pathogenicity island is predominately found in E. coli strains of the phylogenetic group B2 and in some other species of Enterobacteriaceae (1, 2). The pks island carries the genes clbA to clbS, which encode modular nonribosomal peptide synthetases (NRPSs), polyketide synthases (PKSs), and accessory enzymes (1). This complex biosynthetic assembly line is responsible for the synthesis of colibactin, which belongs to the chemical family of hybrid polyketide/nonribosomal peptide (PK-NRP) compounds. Mature colibactin has a complex structure of highly unstable and reactive molecules forming DNA adducts (3–12). However, the complete structure of genotoxic colibactin is not yet fully elucidated.

Colibactin is genotoxic for eukaryotic (1) as well as prokaryotic cells when the ClbS resistance protein is not produced (13). Colibactin-induced DNA damages result from


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the formation of interstrand DNA cross-links leading to DNA double-strand breaks (12, 14). These DNA damages can lead to gene mutations, chromosomal instability, and senescence (15, 16), and in various mouse models colibactin-producing E. coli strains promote intestinal tumor progression (17–20).

Epidemiological studies show a high prevalence of pks-carrying E. coli strains in biopsy specimens from colorectal cancer patients (17, 21, 22). However, the presence of a pks+ strain in gut microbiota is not sufficient to induce colorectal cancer. Other factors have been shown to promote cell transformation and/or to potentiate the bacterial genotoxicity of these bacteria. For example, deoxynivalenol, a food contaminant, exacerbates the genotoxic effect linked to colibactin in animals colonized by pks+ E. coli (23). Environmental factors such as iron concentration can also directly regulate colibactin production (24). However, inflammation is a more significant factor, which seems to be required in colibactin-associated carcinogenesis (17, 20). Inflammation itself is critical for tumor progression (by promoting cell proliferation, survival, and migration) but it also drives modifications in microbiota composition and the expansion of E. coli (25). It has been shown that a proinflammatory cancer microenvironment could increase the expression of clb genes, at least at a transcriptional level (26). Recently, an association between colibactin-producing E. coli and enterotoxigenic Bacteroides fragilis, another procarcinogenic bacterial species, was also noted on the colonic mucosa of patients with familial adenomatous polyposis (FAP), who are highly susceptible to colorectal cancer (20). A synergy between these two bacterial species was observed in tumor formation in an FAP murine model (20).

Polyamines are essential for cell proliferation and have been shown to play a crucial role in carcinogenesis (27, 28). They are small aliphatic molecules involved in protein synthesis and regulation, DNA integrity, stress resistance. (29). Infections by microorganisms and chronic inflammation can interfere with polyamine catabolism and increase the formation of damaging oxidative compounds, contributing in fine to carcinogenesis (30, 31). Polyamines are found at high concentrations in colorectal cancer tissues even at precancerous states (32) but are also produced by intestinal bacteria such as E. coli (33).

In this study, we investigated the putative roles of polyamines in colibactin genotoxic activity. We demonstrate that spermidine is involved in colibactin synthesis and the associated genotoxicity.

RESULTS

The spermidine biosynthetic pathway is required for full genotoxicity of colibactin-producing E. coli. To test the impact of the endogenous spermidine-putrescine pathway on colibactin-producing E. coli genotoxic activity, mutants inactivated for the speB, speC, speE, and speG genes (Fig. 1A) were engineered in E. coli strain DH10B, which harbors the pks island on a bacterial artificial chromosome (Table 1) (34). The production of colibactin by each mutant was first monitored through bacterium-host cell interactions and subsequent observation of the formation of large senescent cells (megalocytosis) (Fig. 1). Inactivation of speB, speC, and speE genes but not of speG resulted in a decrease in the megalocytosis effect (Fig. 1A), with no effect on bacterial growth (data not shown). Therefore, spermidine biosynthesis, and especially the final step catalyzed by the spermidine synthase SpeE, has more impact on colibactin genotoxic activity than putrescine biosynthesis. We confirmed the loss of the megalocytosis phenotype in the ΔspeE mutant and its restoration after plasmid complementation (Fig. 1B). The inactivation of speE in pks+ E. coli from other genetic backgrounds (i.e., the E. coli M1/S commensal strain and the E. coli SP15 pathogenic strain isolated from neonatal meningitis) also resulted in a decrease in the megalocytosis effect (Table 1 and Fig. 1C). We then measured the genotoxic activity of the bacteria by quantifying histone H2AX phosphorylation in response to DNA damage in the infected cells (1, 15, 35). We confirmed that the inactivation of the speE gene in the DH10B pks+ strain markedly decreased the genotoxic effect induced by colibactin (Fig. 2), whereas the complemented mutant was fully genotoxic (Fig. 2).
Spermidine is required for full genotoxicity of colibactin-producing *E. coli*. In order to determine whether the decreased genotoxicity of the ΔspeE mutant is associated with the spermidine synthase SpeE or to spermidine itself, we added increasing concentrations of spermidine in the interaction medium during HeLa cell infection.

**FIG 1** Endogenous spermidine biosynthesis is involved in the *E. coli* colibactin-associated megalocytosis phenotype. (A) Presentation of the polyamine biosynthetic pathways (34). DH10B pks*+/H11001 mutants with mutations of genes speB, speC, speE, and speG were tested for the megalocytosis phenotype in infected HeLa cells, as previously described (1). The phenotypes of HeLa cells resulting from infection with the different mutants are shown. (B and C) Cytotoxic effects of colibactin produced by *E. coli* strains DH10B pks*+/H11001, M1/5, and SP15 and their derivatives were determined by quantification of megalocytosis. At the end of HeLa cell infection, the methylene blue protein staining was quantified by measurement of absorbance at the optical density at 660 nm. The multiplicity of infection (MOI) was 200. Data were pooled from three independent experiments. ***, *P* < 0.001, and **, *P* < 0.01, by 1-way analysis of variance (ANOVA). All bar graphs show mean values ± standard errors of the mean (SEM). NS, not significant.

**TABLE 1** *E. coli* strains and plasmids used in this study

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Genotype or phenotype</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>E. coli</em> strains</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DH10B pks*+</td>
<td>K-12 laboratory strain carrying pBACpks, Cm'</td>
<td>1</td>
</tr>
<tr>
<td>DH10B pks* ΔspeB</td>
<td>speB mutant of strain DH10B pBACpks, Cm' Kan'</td>
<td>This study</td>
</tr>
<tr>
<td>DH10B pks* ΔspeC</td>
<td>speC mutant of strain DH10B pBACpks, Cm' Kan'</td>
<td>This study</td>
</tr>
<tr>
<td>DH10B pks* ΔspeE</td>
<td>speE mutant of strain DH10B pBACpks, Cm' Kan'</td>
<td>This study</td>
</tr>
<tr>
<td>DH10B pks* ΔspeG</td>
<td>speG mutant of strain DH10B pBACpks, Cm' Kan'</td>
<td>This study</td>
</tr>
<tr>
<td>DH10B pks* ΔspeE + p-speE</td>
<td>DH10B pks* ΔspeE carrying p-speE, Cm' Kan' Amp'</td>
<td>This study</td>
</tr>
<tr>
<td>DH10B pks* ΔclbA</td>
<td>clbA mutant of strain DH10B pBACpks, Cm'</td>
<td>13</td>
</tr>
<tr>
<td>DH10B pks* ΔclbS</td>
<td>clbS mutant of strain DH10B pBACpks, Cm'</td>
<td>13</td>
</tr>
<tr>
<td>M1/5 ΔspeE</td>
<td>speE clbS double mutant of strain DH10B pBACpks, Cm'</td>
<td>This study</td>
</tr>
<tr>
<td>M1/5 ΔspeE</td>
<td>speE mutant of strain M1/5, Kan'</td>
<td>This study</td>
</tr>
<tr>
<td>SP15 ΔspeE</td>
<td>Extrainestinal pathogenic <em>E. coli</em> strain isolated from spinal fluid of neonate with meningitis, O18K1 serotype, colibactin genotoxin producer</td>
<td>50</td>
</tr>
<tr>
<td>SP15 ΔspeG</td>
<td>speG mutant of strain SP15, Kan'</td>
<td>This study</td>
</tr>
<tr>
<td>SP15 ΔspeE</td>
<td>speE mutant of strain SP15, Kan'</td>
<td>This study</td>
</tr>
<tr>
<td>Plasmid p-speE</td>
<td>pSC-A plasmid carrying wild-type speE gene, Amp' Kan'</td>
<td>This study</td>
</tr>
</tbody>
</table>
The production of colibactin was again measured by quantifying histone H2AX phosphorylation and megalocytosis assay (see Fig. S1 in the supplemental material). Spermidine supplementation of the ΔspeE mutant restored its genotoxicity in a dose-dependent manner (Fig. 3). The spermidine acetyltransferase SpeG catalyzes spermidine acetylation into physiologically inert N-acetylspermidine. While screening the impact of the putrescine-spermidine pathway on colibactin-associated megalocytosis phenotype, we observed no ΔspeG mutant defect (Fig. 1A). We hypothesized that by abolishing SpeG activity, spermidine would accumulate in the bacteria and boost the genotoxic activity of pks\textsuperscript{+} E. coli. To test this hypothesis, we quantified H2AX phosphorylation in response to DNA damages after HeLa cell infection by E. coli SP15 and deletion mutants of speE and speG genes (Fig. 4). Compared to the wild-type strain, we observed a decrease in the genotoxic activity of SP15 ΔspeE (Fig. 4). However, SP15 ΔspeG with impaired catabolism of spermidine induced 20% more DNA damage than the wild-type SP15 strain (Fig. 4). These results confirmed that spermidine is the key player in the interaction between the putrescine-spermidine pathway and colibactin-associated genotoxic ac-
tivity. However, a ΔpotD spermidine import mutant of E. coli strain SP15 was not impaired in its genotoxicity (see Fig. S2 in the supplemental material), suggesting that spermidine endogenous production is sufficient to support colibactin production in this in vitro assay. Furthermore, we tested other polyamines and showed that spermine and norspermidine could restore the genotoxic activity in a ΔspeE mutant (see Fig. S3 in the supplemental material).

Spermidine is directly involved in colibactin biosynthesis. To determine the level at which spermidine acts, we quantified the autotoxicity linked to colibactin production directly in bacteria. In fact, DNA damage can occur in toxigenic bacteria mutated for the ClbS resistance protein, leading to activation of the SOS response and then decreased growth (13). We constructed a ΔspeE ΔclbS double mutant in the E. coli strain DH10B pks/H11001 (Table 1) and compared its growth after 17 h in LB broth to ΔclbS and ΔspeE mutants. As expected, there was a decrease in the number of ΔclbS mutant CFU compared to the wild-type strain (Fig. 5). Reduced CFU counts were also observed for the ΔspeE ΔclbS double mutant compared to the ΔspeE mutant, but to a lesser extent than for the ΔclbS mutant (Fig. 5). Therefore, speE mutation significantly decreases colibactin autotoxicity in a ΔclbS mutant, suggesting decreased production of genotoxin by the bacteria when the polyamine pathway is disrupted (Fig. 5).

To confirm the involvement of spermidine in colibactin production, we monitored the DNA cross-linking activity of colibactin-producing E. coli in exogenous DNA, in an acellular assay. Following incubation with bacteria, plasmid DNA was purified and analyzed by electrophoresis on agarose gel under denaturing conditions to highlight DNA interstrand cross-linking with delayed migration compared to linearized single-stranded DNA (Fig. 6) (14). This experiment showed that speE mutation greatly atten-
uates DNA cross-linking activity, which was observed only for the highest bacterial dose (Fig. 6). The ability of the ΔspeE mutant to induce DNA cross-links was restored by either exogenous addition of spermidine or speE complementation (Fig. 6) or transcomplementation by polyamine production of a wild-type E. coli strain without pks island in coculture (see Fig. S4 in the supplemental material). These experiments performed without eukaryotic host cells suggest that spermidine has a direct role in the production of colibactin, which results in fully genotoxic bacteria.

Considering that mature colibactin is not yet directly quantifiable, we took advantage of the stability of the N-myristoyl-D-asparagine moiety cleaved by the ClbP peptidase in the late activation step of inactive precolibactin (3, 4) to indirectly access the amount of colibactin produced, using LC-MS (36) (Fig. 7). Quantification of N-myristoyl-D-asparagine in culture supernatants revealed that the amount of colibactin produg motif was drastically decreased in the ΔspeE mutant compared to the E. coli DH10B pks⁻ strain, and partially restored by complementation with p-speE (Fig. 7).

**FIG 5** Deletion of the spermidine synthase SpeE decreases colibactin autotoxicity in a ΔclbS mutant. Enumeration of culturable bacterial cells in the stationary growth phase. The bacteria were pregrown in LB to reach an exponential growth (OD₆₀₀ = 0.4). A total of 2 × 10⁶ bacteria/ml were then inoculated in LB and grown for 17 h before being plated on LB agar plates to determine CFU. The median and individual results of four independent experiments are shown. ***, P < 0.001, and **, P < 0.01, by 1-way ANOVA. NS, not significant.

**FIG 6** Spermidine is involved in DNA cross-linking activity. DNA cross-linking was observed by cultivating the E. coli strains with linearized plasmid DNA. Two hundred nanograms of linearized pUC19 plasmid was added to 1 × 10⁶, 2 × 10⁶, or 4 × 10⁶ bacteria/well of wild-type (WT) E. coli strain DH10B pks⁻, the ΔclbA and ΔspeE mutants, and the complemented derivative. When indicated, 20 μg/ml of spermidine was added to the interaction medium during infection. DNA was then purified, loaded on agarose gel, and run under alkaline denaturing conditions. DNA with covalent interstrand cross-links is nondenaturable and displays delayed migration compared to denatured single-stranded DNA (lower band). This image is representative of three independent experiments.
Supplementation of spermidine during growth increased the production of N-myristoyl-D-asparagine by the ΔspeE mutant (Fig. 7).

We tested whether spermidine altered clb gene expression by using clb reporter strains previously designed in E. coli Nissle 1917 (37) (see Fig. S5 and Text S1 in the supplemental material). We observed modifications of expression after both ΔspeE mutation and spermidine supplementation, suggesting that spermidine can modify expression of pks genes.

**DISCUSSION**

Our work demonstrates the role of spermidine in the production of the genotoxin colibactin. We demonstrate that the endogenous spermidine biosynthesis pathway is involved in colibactin-associated genotoxicity. As a ΔspeE mutant is impaired in its genotoxic activity in acellular DNA cross-linking assay, a mechanism mediated by the targeted eukaryotic cell (such as invasion) was ruled out. Spermidine is not necessary, however, as traces of colibactin-related toxicity remain after speE mutation, as demonstrated both by autotoxicity in the ΔspeE ΔclbS double mutant and DNA cross-link formation at the highest bacterial concentration. Furthermore, N-myristoyl-D-asparagine, a metabolite directly linked to colibactin synthesis, also undergoes quantitative alteration when the spermidine pathway is inactivated, showing a direct involvement of spermidine in production of the toxin. Not only exogenous spermidine but also other polyamines such as spermine and norspermidine can restore colibactin biosynthesis in ΔspeE mutants. Noteworthy, the three polyamines that cause complementation share structural characteristics: i.e., 3 or more amine functions, 2 of which are free at the ends. Proposed colibactin and precolibactin structures do not contain any spermidine or spermidine-derived motif (3–12). Thus, polyamines may likely be involved in colibactin synthesis with regard to monitoring its regulation. Precisely we have observed changes in the expression of different clb genes related to speE mutation or spermidine supplementation. Although the precise molecular mechanism remains elusive, polyamines are known to be major regulators in E. coli gene expression, especially at the translation step (38). Indeed, enhancement of mRNA and ribosome interaction leads to increased translation of proteins that belong to the “polyamine modulon” as it is called by the Igarashi and Kashiwagi team (38). Alternatively, polyamines could also act through facilitating transport or export of the genotoxin.
Spermidine and polyamines have been described as important players in different host-pathogen relationships. In the plague agent *Yersinia pestis*, for instance, an intact putrescine-spermidine pathway is essential to achieve biofilm formation, which is in turn important for transmission by fleas (39). A higher virulence linked to an increase in the spermidine content exists in *Shigella* sp., a bacterium responsible for dysentery and closely related to *E. coli* (40). During the pathoadaptive evolutionary changes in *E. coli* that led to this pathogen, the speG gene became defective due to diverse mutations. Spermidine accumulation in bacteria is a key factor for the invasion of *Shigella* sp. and entero-invasive *E. coli* (EIEC), which share the same invasive process (40, 41). The production and response of colicin E7, an *E. coli* bacteriocin, are also regulated by polyamines (42). Spermidine and putrescine are indeed important for triggering toxin production after DNA damage because polyamines are mediators of the SOS response that regulates colicin E7. Exogenous spermidine decreases the susceptibility of *E. coli* to colicin E7, which can be attributed to a downregulation of the transporters of the toxin by the polyamine (42).

Spermidine is highly linked to cancer, especially colorectal cancer. Polyamines are associated with cell proliferation. Therefore, host cell polyamine content is upregulated in colorectal cancer tissues. It relies on both an increase in polyamine production and impaired catabolism (28). The host is not the only source of polyamine in the digestive tract. Bacteria in gut microbiota also produce polyamines and are one of the main sources of these compounds in the lower intestine (43). The diversity of the microorganisms makes the metabolism and composition of the polyamine pool diverse (44, 45), and it can be even more complex as a result of collective pathways within microorganisms as well as polyamines produced by eukaryotic cells (46). Food is another source of polyamines in the gut, especially nutrients such as fruits, wheat germ, cheeses, mushrooms, and nuts. (47). In our study, we demonstrated that exogenous spermidine produced by another *E. coli* strain could support colibactin biosynthesis by a spermidine mutant. This strongly suggests that in the gut, where spermidine is abundant, colibactin-producing bacteria can take advantage of this metabolite, whatever the origin of its production (microbiota, host, or food). Recently, the external supply of polyamine spermidine was also suggested to extend life span and to provide neuroprotective and cardioprotective properties, for example (48). To date, the literature on these beneficial effects of spermidine has been limited to model organisms and to epidemiological studies that link dietary uptake with longevity or pathology. Clinical trials aimed at increasing the uptake of this polyamine or even the administration of probiotics that enhance the microbial synthesis appear feasible (48). However, whether spermidine supplementation in humans would increase tumor growth or modify microbiota remains unknown and should be addressed in light of the fact that a polyamine-enriched environment could promote and may even trigger the production of the genotoxin colibactin. This reinforces the hypothesis of deleterious synergy between microbiota and host cell metabolism during carcinogenesis.

MATERIALS AND METHODS

**Bacterial strains, mutagenesis, and growth conditions.** The bacterial strains used in this study are listed in Table 1 and Table S1 in the supplemental material (Text S1). For genetic manipulations, *E. coli* strains were routinely grown at 37°C under shaking in 5 ml of lysogeny broth (LB Lennox; Invitrogen). Appropriate antibiotics were added to the medium when required (chloramphenicol, 25 μg/ml; kanamycin, 50 μg/ml; carbenicillin, 50 μg/ml). Inactivation of the genes speB, speC, speE, and speG was performed by using the lambda Red recombinase method (49) with pairs of primers presented in Table 2 and Table S2 in the supplemental material. Allelic exchanges were confirmed by PCR. For complementation, the speG gene was PCR amplified using the primers speE CompF and speE CompR cloned into pSC-A-amp/kan using the StrataClone PCR kit (Agilent) (see p-speE in Table 1).

For the megalocytosis assay, genotoxicity quantification, DNA cross-linking assay, and N-myristoyl-o-asparagine quantification, *E. coli* strains were pregrown overnight at 37°C with shaking in Dulbecco’s modified Eagle’s medium (DMEM)-HEPES (Gibco), a polyamine-free medium. Overnight cultures were then diluted 1:50 in DMEM-HEPES and grown until they reached an optical density at 600 nm (OD600) of 0.6 and then processed for experiments.
TABLE 2 Primers used in this study

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
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</thead>
<tbody>
<tr>
<td>speB_P1</td>
<td>TACTGGCGTTGCGGTATATGGGACCTTCTGGGTGTGGTGGTAGTGGTAGGAGCTGCTTTC</td>
</tr>
<tr>
<td>speB_P2</td>
<td>TAATAGCGGCGATGAGGTCAGGCACCGCAATCATCCTGGCTCTGGTAGCTAGATATCTCTTAG</td>
</tr>
<tr>
<td>speB_F</td>
<td>GGTTTACCGGCGGTCCATCG</td>
</tr>
<tr>
<td>speC_F</td>
<td>CCCTATCCCTATCATATTCTATCTCC</td>
</tr>
<tr>
<td>speP1</td>
<td>GTTGGTATATACCGGATTCCTAGCCCGAGAGCATATGATATCTCTTAG</td>
</tr>
<tr>
<td>speP2</td>
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<td>speR</td>
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</tr>
<tr>
<td>speCompR</td>
<td>ATCTCGAGGTTTTGGCTGGTAGCAGATATCG</td>
</tr>
</tbody>
</table>

For the autotoxicity assay, E. coli strains and derivatives were pregrown in LB to reach an exponential growth (OD_{560} = 0.4). A total of 2 × 10^8 bacteria/ml were then inoculated in LB and grown for 17 h before being plated on LB agar plates to determine CFU counts.

Determining the megacytosis and genotoxic effect induced by colibactin. The megalocytosis and genotoxic effect induced by colibactin were determined as previously described (35). Briefly, HeLa cells were dispensed in a 96-well cell culture plate (5 × 10^3 to 7.5 × 10^3 cells/well) and incubated for 24 h. For bacterial infections, cell cultures were infected with a multiplicity of infection (number of bacteria per HeLa cell at the onset of infection) ranging from 100 to 400. Four hours postinoculation, cells were washed three times with Hanks balanced salt solution (HBSS) and incubated in cell culture medium with 200 μg/ml gentamicin until analysis.

For megalocytosis quantification, cells were incubated for 72 h before protein staining with methylene blue (1% wt/vol in 0.01 M Tris-HCl [pH 8.5]). The methylene blue was extracted with 01 N HCl. The quantification of staining was measured at OD_{660}.

For H2AX phosphorylation quantification, cells were incubated for 3 h overnight before fixation with formaldehyde, permeabilization, and blocking, as previously described (35). Cells were then incubated for 2 h at room temperature with rabbit monoclonal anti-γH2AX antibody (1:200 [Cell Signaling Technology]). An infrared fluorescent IRDye™800CW-conjugated goat anti-rabbit secondary antibody was added to the cells, and DNA was counterstained with RedDot2 (1:500 [Biotium]). DNA and γH2AX were visualized simultaneously using an Odyssey Infrared Imaging Scanner (Li-Cor Biosciences) with 680- and 800-nm channels. Relative fluorescence units for γH2AX per cell (as determined by γH2AX divided by DNA content) were divided by untreated controls. Results were then divided by the mean ratio obtained for the wild-type strain after infection with the same infectious dose to determine percentage change in phosphorylation of H2AX levels relative to this reference strain.

DNA cross-linking assay. The assay was performed as previously described (14). Briefly, linearized DNA was obtained by digesting pUC19 plasmid with BamHI (NEB). Purified linearized DNA was quantified and diluted to obtain a 200-ng/μl stock solution. For bacterium-DNA interactions, 1 × 10^6 to 6 × 10^6 CFU were cultured with linearized DNA for 4 h at 37°C without shaking. Following centrifugation for 5 min at 5,000 × g to pellet bacteria, the DNA present in the supernatants was purified using the PCR purification kit (Qiagen) according to the manufacturer’s recommendations.

Denaturing agarose gel was prepared by dissolving 1.0 g of agarose in 100 ml of a 100 mM NaCl and 2 mM EDTA solution (pH 8.0). The gel was then soaked (2 h) in an alkaline running buffer solution (40 mM NaOH and 1 mM EDTA [pH ~12.0]). One hundred nanograms of each DNA sample was loaded on the agarose gel. The gel was run for 45 min at 1 V/cm and then 2 h at 2 V/cm. The gel was then neutralized for a total of 45 min in a 100 mM Tris (pH 7.4) buffer solution containing 150 mM NaCl, and the neutralizing solution was refreshed every 15 min. The gel was stained with GelRed for 20 min and revealed with UV exposure using the ChemiDoc imaging system (Bio-Rad).

N-Myristoyl-α-asparagine (colibactin prodrug motif) quantification by liquid chromatography-mass spectrometry. The colibactin prodrug motif was quantified as previously described (36). Briefly, precultivated strains were grown in DMEM-HEPES at 37°C for 18 h under shaking (240 rpm). Supernatants of cultures were obtained by centrifugation of bacterial cells at 3,200 × g for 15 min and were filtered on 0.2-μm-pore membranes. Each strain was cultured in triplicate (derived from three independent clones), and each supernatant was analyzed by liquid chromatography-tandem mass spectrometry (LC-MS/MS).

Quantification experiments were performed with ultraperformance liquid chromatography high-resolution/heated electrospray ionization mass spectrometry (UPLC-HR/ESI-MS). The data were recorded on a Thermo Scientific Q Exactive Exactive hybrid quadrupole-Orbitrap mass spectrometer coupled to a Dionex Ultimate 3000 UPLC. The following solvent gradient (A = H_2O + 0.1% formic acid, B = acetonitrile + 0.1% formic acid with B at 30% from 0 to 1 min, 30 to 95% from 1 to 6 min, and 95% from 6 to 7 min at a flow rate of 0.5 ml/min) was used on a Phenomenex Kinetex 5-μm EVO C_{18} (50- by 2.1-mm) column at 30°C. The mass spectrometer was operated in positive-ionization mode at a scan range of 200-
to 500 mV and a resolution of 35,000. The spray voltage was set to 3.5 kV, the S-lens to 35, and the auxiliary gas heater temperature to 438°C, and the capillary temperature to 270°C. Absolute quantification was achieved by using a Schotten-Baumann reaction-derived N-myristoyl-D-asparagine (isomer of the N-myristoyl-L-asparagine colibactin cleavage product) as a standard. Data were obtained from undiluted cell-free sample supernatants and analyzed for N-myristoyl-D-asparagine, and concentrations were calculated using Thermo Xcalibur 2.2 Quan Browser.

**Statistical analysis.** Statistical analyses were conducted using GraphPad Prism 6.01. The mean and the standard error of the mean (SEM) are shown in the figures, unless otherwise stated. P values were calculated by a one-way analysis of variance (ANOVA) followed by a Bonferroni post hoc test. A P value of <0.05 was considered statistically significant.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mSphere.00414-19.

TEXT S1, DOCX file, 0.1 MB.

FIG S1, TIF file, 0.6 MB.

FIG S2, TIF file, 1.3 MB.

FIG S3, TIF file, 0.7 MB.

FIG S4, EPS file, 0.9 MB.

FIG S5, TIF file, 1.1 MB.

TABLE S1, DOCX file, 0.1 MB.

TABLE S2, DOCX file, 0.1 MB.

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**REFERENCES**


Colibactin Synthesis Is Linked to Polyamine Metabolism


