Competence-Independent Activity of Pneumococcal Enda Mediates Degradation of Extracellular DNA and Nets and Is Important for Virulence

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Abstract
Membrane surface localized endonuclease Enda of the pulmonary pathogen Streptococcus pneumoniae (pneumococcus) is required for both genetic transformation and virulence. Pneumococcus expresses Enda during growth. However, it has been reported that Enda has no access to external DNA when pneumococcal cells are not competent for genetic transformation, and thus, unable to degrade extracellular DNA. Here, by using both biochemical and genetic methods, we demonstrate the existence of Enda-mediated nucleolytic activity independent of the competence state of pneumococcal cells. Pneumococcal mutants that are genetically deficient in competence development and genetic transformation have extracellular nuclease activity comparable to their parental wild type, including their ability to degrade neutrophil extracellular traps (NETs). The autolysis deficient ΔlytA mutant and its isogenic choline-treated parental wild-type strain D39 degrade extracellular DNA readily, suggesting that partial cell autolysis is not required for DNA degradation. We show that Enda molecules are secreted into the culture medium during the growth of pneumococcal cells, and contribute substantially to competence-independent nucleolytic activity. The competence-independent activity of Enda is responsible for the rapid degradation of DNA and NETs, and is required for the full virulence of Streptococcus pneumoniae during lung infection.

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Introduction
Enda is reported to be a membrane-localized pneumococcal endonuclease [1]. It was first implicated to play a role in genetic transformation by Kohoutova [2], and subsequently confirmed by Lacks and colleagues [3–5]. During genetic transformation, Enda degrades one strand of double stranded DNA (dsDNA) and converts it into single stranded DNA (ssDNA) for uptake and recombination [6]. Acid soluble DNA fragments or nucleotides generated during DNA degradation are released into the culture medium [4], and are only detectable during competence development [7]. These observations were further confirmed by Berge and colleagues [8]. They showed that the ΔcomD mutant, which lacks the histidine kinase receptor required for the induction of the competence regulon and genetic transformation, is both unable to develop competence and degrade DNA. In addition, the comEd and cglABCDGF operons, which encode the apparatus for DNA binding and uptake, are required for Enda-mediated DNA degradation. These authors proposed that DNA attaches to the apparatus in competent pneumococcal cells, allowing the membrane-localized Enda to gain access to the extracellular DNA. For pneumococcal cells not under the competent state, Enda is unable to gain access to donor DNA, and is incapable of degrading extracellular DNA.

Demonstration of Enda’s role in pneumococcal virulence is comparatively recent. An Enda-deficient mutant was identified in a signature-tagged mutagenesis screen in mouse lungs [9]. Furthermore, Enda degrades neutrophil extracellular traps (NETs) [10]. Composed of both DNA and antibacterial proteins, NETs are released by activated neutrophils to capture and kill bacterial cells [11]. Thus, it is conceivable that rapid degradation of NETs by Enda releases captured pneumococcal cells and facilitates their dissemination.

In this study, we present experimental evidence that majority of the nucleolytic activities of Enda are independent of competence development, and the “competence-independent activity” of Enda contributes to lung infection. In addition, Enda is secreted during pneumococcal growth.

Materials and Methods
Synthetic CSP1, Bacterial Strains and Growth Conditions
CSP1 (≥95% purity) was synthesized by Elim Biopharm. S. pneumoniae strains are listed on Table 1. Wild-type strain D39 [12] was a gift from Dr. David Briles (University of Alabama-Birmingham). R6 is a capsule-deficient mutant derived from D39 [13] whereas 0100993 is a highly encapsulated serotype III clinical isolate [14]. Mutant strains ΔcomD, ΔcomA, ΔcglABCDGF,
ΔlytA, and ΔendA were generated by nonpolar deletions in D39 using the Janus cassette as previously described [15]. Strain JC0923, which carries both an insertion of the lacZ gene under the control of the comX promoter and a nonpolar deletion in the comA gene, was generated by transforming ΔcomA DNA from D39pcbpD::lacZ [16,17]. Bacteria were streaked from frozen stocks onto THB (Todd Hewitt Broth) agar containing 5% defibrinated horse blood at 37°C with 5% CO₂. Fresh colonies were grown in THB or CTM (complete transformation medium) [18] to desired density.

**DNA Degradation Assay**

Pneumococcal cells were grown in THB to early log phase around 10⁸ CFU/ml (1 ml). The OD 600 nm at this bacterial concentration for D39, R6 and 0100993 were 0.15, 0.1 and 0.4, respectively. The differing OD600nm at 108 CFU/ml between strains D39 and 0100993 is due to the presence of different pneumococcal strains in CTM to 108 CFU/ml (1 ml). The OD 600 nm at this bacterial concentration for D39pcbpD::lacZ [16,17]. Bacteria were streaked from frozen colonies on THB agar incorporated with 166 μg/ml salmon sperm DNA. After 12 hr, DNA plates were flooded with 5 ml of 1 N HCl as previously described [19]. DNA degradation was inhibited by adding 200 ml of fresh THB and added to column purified PCR products of the streptomycin resistance rpsL gene (30 μg). At indicated time points, a 100 μl aliquot of the bacteria-DNA mixture was withdrawn and immediately centrifuged at 12000 rpm for 2 min. DNA-containing supernatants were stored at −80°C to prevent further degradation. The integrity of DNA was visualized by agarose gel electrophoresis. For degradation of DNA in the CTM medium, pneumococcal strains were grown in THB to 10⁶ CFU/ml (1 ml), washed, and resuspended in 1 ml of CTM. DNA degradation appeared as transparent zones of clearance.

**Degradation of [α32P]-dATP-labeled DNA**

P-32 labeled donor DNA was amplified by PCR using 1x PCR master mix (Thermoscientific) and D39 chromosomal DNA (100 μl reaction size). PCR was carried out with the rpsL primers in the presence [α32P]-dATP (Amersham; specific activity 111×10⁶ Bq mmol−1) for 40 cycles with the following settings: 30 seconds at 94°C, 30 seconds at 54°C, 60 seconds at 72°C, followed by a 10 min extension step. Hot PCR products were purified by QiAquick PCR Purification Kit (Qiagen). Hot DNA (5 μg) was exposed to 10⁸ CFU/ml ΔcomA cells (1 ml) with or without CSP1 stimulation. The ΔendA cells (10⁸ CFU/ml) and THB were used as controls. To determine the extent of DNA degradation, an aliquot of 100 μl was withdrawn from each group at indicated time points after incubation. The samples were centrifuged at 12000 rpm for 1 min to remove bacterial cells.

### Table 1. *S. pneumoniae* strains used in this study.

| Strains | Relevant characteristics | Reference |
|---------|--------------------------|-----------|
| D39 | Wild-type | [12] |
| AD2064 (ΔcomCDE) | A competent deficient D39 derivative generated by deleting the comCDE operon using the Janus cassette | This work |
| AD1863 (ΔcglABCDEFG) | A transformation deficient D39 derivative generated by deleting the cglABCDEFG operon using the Janus cassette | This work |
| AD1737 (ΔlytA) | A D39 derivative deficient in major autolysin generated by deleting the lytA gene using the Janus cassette | This work |
| AD1762 (ΔendA) | A transformation deficient D39 derivative generated by deleting the endA gene using the Janus cassette | This work |
| AD0049 (ΔcomA) | A D39 derivative deficient in spontaneous competence generated by deleting the comA gene using the Janus cassette | This work |
| IN1643 (D39pcmx::lacZ) | D39 with a promoterless lacZ reporter gene fused behind the comX promoter | This work |
| JC0923 | D39comX::lacZ strain with a deletion in the comX gene | This work |
| CPM3 | R6 with a promoterless lacZ reporter gene fused behind the comX promoter | [31] |
| CP1296 | R6 strain with a modified rpsL gene that confers resistance to streptomycin | [15] |
| R6 | A capsule-deficient derivative of D39 | This work |
| AR2042 (R6ΔcomD) | A competence-deficient R6 derivative generated by deletion of the comD gene using the Janus cassette | This work |
| AR1779 (R6ΔendA) | A R6 derivative generated by the deletion of the endA gene using the Janus cassette | This work |
| TIGR4 | Wild-type | [35] |
| AT2236 (TIGR4ΔcomD) | A competence-deficient TIGR4 derivative generated by deleting the comD gene using the Janus cassette | This work |
| AT1762 (TIGR4ΔendA) | A transformation-deficient TIGR4 derivative generated by deleting the endA gene using the Janus cassette | This work |
| 0100993 | Serotype 3 clinical isolate | [14] |
| ST2012 (0100993Δcap3A) | A capsule-deficient 0100993 derivative generated by deleting the cap3A gene using the Janus cassette | This work |
| AG2236 (0100993ΔcomD) | A competence-deficient 0100993 derivative generated by deleting the comD gene using the Janus cassette | This work |
| AG1762 (0100993ΔendA) | A transformation-deficient 0100993 derivative generated by deleting the endA gene using the Janus cassette | This work |

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Large intact hot DNA fragments within the supernatant were removed by QIAquick PCR Purification columns. The filtrates (20 µl), which contained the hot nucleotides and small DNA fragments (<100 bp), were spotted on filter paper, dried, and exposed in a phosphoimager exposure cassette for 8 hr and scanned with the FujiFilm FLA-3000 PhosphorImager. Phosphoimager signals were quantified by Image J. For quantitative determination of released nucleotides/small DNA fragments, the filtrates (20 µl), which contained the hot nucleotides and small DNA fragments (<100 bp), were mixed thoroughly with 5 ml of Cytoscint® Scintillation Cocktail (Thermoscientific). Radioactivity was quantified by using a Tri-Carb® 2100TR Liquid Scintillation Counter (Perkin Elmer).

**Analysis of the EndA Secretion**

Pneumococcal strains TIGR4, 0100993, and D39 and its derivatives ΔcomD and ΔlytA, were cultured in THB to ~ 10^9 CFU/ml. One ml of culture was withdrawn and subjected to centrifugation. Bacterial cells were then washed three times with PBS to remove cultural medium and resuspended into 1 ml of fresh THB. Resuspended bacterial strains were allowed to grow in the 37°C incubator supplemented with 5% CO₂. At 0, 30, 60 and 90 min, supernatant samples of each bacterial culture were
collected. Pneumococcal cells were removed using the 0.25 μm filters. 20 μl of bacterial cells or supernatant was boiled with 5 μl of 5x SDS loading buffer (0.25% Bromophenol blue, 0.5 M dithiothreitol, 10% Glycerol, 0.25 M Tris-Cl, pH 6.8) for 5 min, and subjected to SDS-PAGE in an acrylamide gel incorporated with 15 μg/ml of salmon sperm DNA. Following electrophoresis, the gel was washed with pure water to remove SDS, allowing renaturation of proteins. After 12 hr of incubation in 2 mM MgCl2 solution at 37°C, the gel was stained with 10 μg/ml of ethidium bromide to visualize bands of DNA clearance caused by pneumococcal nuclease. A band of clearance at 25 kDa indicates the presence of EndA nucleolytic activities.

To further compare the amount of EndA secreted by pneumococcus growing in THB versus in CTM, washed D39 cells (108 CFU/ml) were resuspended into 1 ml of fresh THB or CTM, respectively, allowed to grow in the 37°C incubator supplemented with 5% CO2 for 2 hr. One ml of cell-free supernatant from D39 cells growing in THB were diluted 10, 100 and 1000 folds. Nuclease activity of different diluents was compared against cell-free supernatant from D39 cells grown in CTM. Each supernatant sample was incubated with hot PCR product at 37°C for 1 hr. The release of nucleotides/small DNA fragments was quantified by using the Tri-Carb® 2100TR Liquid Scintillation Counter (Perkin Elmer).

Genetic Transformation Assay
Pneumococcal JC0923 cells were grown to OD 600 nm 0.15 in THB (pH 6.8), washed and resuspended in fresh THB (pH 8.3) or CTM, and stimulated with 400 ng/ml CSP1. Donor DNA was added to the final concentration of 30 μg/ml. Donor DNA was generated by amplifying a mutated rpsL gene and its flanking regions from a streptomycin resistant strain CP1296 (Table 1), using the following primers: rpsL upper 5’-GGGCTAGTGGTAGTGGTGGG-3’; rpsL lower 5’-CGGAAGTGCGTGGG-3’.
GAATGCACG-3’ (PCR product size: 1633 bp). The transformation mix was then incubated at 37°C with 5% CO₂ for 2 hr. Transformants were selected on THB agar supplemented with 100 μg/ml streptomycin after serial dilutions.

**Activation Assay of comX and Promoter**

 JC0923 cells were grown in THB (pH 6.8) until OD 600 nm of 0.15, washed and resuspended in fresh THB (pH 8.3) or CTM. CSP1 (400 ng/ml) was added to the bacterial culture and incubated at 37°C for 30 min. β-galactosidase activity was measured according to previously published protocols [20] and expressed as Miller units.

**Neutrophil NETs Degradation Assay**

Human neutrophils (Innovative Research) grown in RPMI medium were exposed to 10 mM H₂O₂ in 37°C for 20 min to induce the formation of NETs [21]. Activated neutrophils were incubated for 60 min with RPMI, 10⁸ cells of D39, ΔcomCDE, ΔeglABCDEFG, or ΔendA, respectively. Neutrophils were fixed with 4% paraformaldehyde and stained with DAPI to visualize NETs degradation by using a fluorescence microscope (Zeiss LSM 700).

**Ethics Statement**

The animal study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The protocol was approved by the Institutional Animal Care and Use Committee (IACUC) at the University of Illinois at Urbana-Champaign (Protocol Number: 12230).

**Mouse Acute Pneumonia Infection**

CD1 mice (6-week old, n = 10) (Charles River) were housed in positively-ventilated microisolator cages with automatic recirculating water, located in a room with laminar, high efficiency particle accumulation-filtered air. The animals received autoclaved food, water, and bedding. Mice were anesthetized with isoflurane and intranasally administered 10⁶ CFU of D39, ΔeglABCDEFG, or ΔendA cells. The infected mice were monitored for 48 hr before the lungs were harvested for bacterial enumeration. Moribund animals that displayed rough hair coat, hunched posture, distended abdomen, lethargy or inability to eat or drink were euthanized. Animal studies were carried out in strict accordance to the protocol (#12230) approved by the IACUC at the University of Illinois at Urbana-Champaign.

**Statistical Analyses**

Statistical analyses of *in vitro* experiments were performed using the Student’s *t*-test and one-way analyses of variance (ANOVA). Statistical significance of bacterial burden in mouse lungs was compared using the GraphPad Prism statistical software package. A significant difference was considered to be *p*<0.05.

**Results**

Degradation of Extracellular DNA by EndA does not Require Components of the Competence Regulon

We compared the ability of the wild-type pneumococcal strain D39 and its isogenic endonuclease-deficient ΔendA mutant to degrade extracellular DNA. DNA was rapidly degraded by D39.
Figure 5. Nucleolytic activity of EndA during competence induction by CSP1. PCR-amplified P-32 labeled donor DNA (5 μg) was exposed to ΔcomA cells in the presence or absence of 400 ng/ml CSP1. The ΔendA cells and THB were used as controls. Experiments were performed in triplicates. (A) Scanned image of small hot DNA fragments (<100 bp) and nucleotides spotted on filter paper and exposed to a phosphomager cassette. (B) Quantification of dots (n = 3) in A. *p < 0.05 when comparing the densitometry number of ΔcomA supplemented with CSP1 against ΔcomA alone at 30 min and 90 min post CSP1 exposure.

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EndA activity that is independent of the competence development in pneumococcus.

DNA Degradation by EndA is not due to Pneumococcal Autolysis

Autolysis of pneumococcus during growth is primarily mediated by the autolysin LytA [24], and may release EndA from cell membrane or cytoplasm into the environment. However, the ΔlytA mutant degraded extracellular DNA as efficiently as D39 [Fig. 1A–B]. In addition, D39 cells treated with 2% choline chloride – which inhibit the autolytic activities of all pneumococcal lytic proteins LytA, LytB, LytC, and CbpD [25–29] – degraded extracellular DNA as efficiently as untreated D39 cells [Fig. 1A–B]. These results indicate that pneumococcal autolysis does not contribute significantly to the competence-independent EndA activity.

Induction of Competence Contributes Minimally to the Overall Amount of DNA Degradation by Pneumococcus

To determine if the extracellular DNA degradation by EndA was dependent on the development of competence, we examined the induction of competence by using the pneumococcal strain JC0923 [Table 1] in the presence or absence of CSP1. JC0923 cells carry a lacZ reporter gene under control of the comX promoter and a deletion of the comA gene. ComX is a competence specific sigma factor that positively regulates the transcription of genes for DNA uptake and recombination [30,31]. ComA is an ABC transporter that exports CSP1 [32]. Therefore, JC0923 serves as an ideal test strain to monitor comX expression and genetic transformation, which could only be triggered by exogenously supplied CSP1. We compared the transformation efficiency of JC0923 in THB and in CTM, a medium that allows high transformation efficiency [18]. As shown in Fig. 2A and 2B, the expression of comX promoter driven LacZ is comparable in both media, suggesting that the
induction of competence by CSP1 in THB was successful. Nevertheless, transformation efficiency in the CTM is 3 times higher than in the THB. JC0923 cells grown in THB were able to degrade the rpsL DNA with equal efficiency in the presence or absence of exogenously supplied CSP1 (Fig. 2C). DNA staining with ethidium bromide over the time course of the experiment showed that the kinetics of DNA degradation was indistinguishable between the JC0923 cells with or without CSP1 treatment (Fig. 2D). Collectively, these results suggest that the contribution of competence induction to the overall DNA degradation by pneumococcus is negligible.

Competence-independent EndA Nucleolytic Activity is Conserved in Different Pneumococcal Strains and is Culture Medium Dependent

To determine if the competence-independent activity of EndA is a widespread phenomenon, we tested the pneumococcal strains R6, 0100993, and TIGR4 as well as their isogenic ΔcomD mutants for their ability to degrade extracellular DNA. D39 was used as positive control for DNA degradation. Pneumococcal strains were grown in THB, washed and resuspended in THB or CTM medium and examined for DNA degradation. As shown in Fig. 3A, R6, 0100993 and TIGR4 possess different nucleolytic activity levels. In contrast, their isogenic ΔendA mutants are unable to degrade DNA. Importantly, the ΔcomD mutants degrade DNA as efficiently as their respective parental strains R6, 0100993 and TIGR4, confirming that the competence system is not important for EndA-mediated DNA degradation. In contrast, their isogenic ΔendA mutants are unable to degrade DNA. Importantly, the ΔcomD mutants degrade DNA as efficiently as their respective parental strains R6, 0100993 and TIGR4, confirming that the competence system is not important for EndA-mediated DNA degradation. In addition, D39 and its unencapsulated derivative R6, as well as 0100993 and TIGR4 possess different nucleolytic activity levels. These results suggest capsule does not influence EndA activity in these strains. However, the nucleolytic activity appears to be influenced by pneumococcal growth. D39 grew at slower rate in the CTM (Fig. 3B). Similarly, TIGR4 and 0100993 also grew slower in CTM (data not shown). Degradation of extracellular DNA was severely impaired when D39 was grown in CTM (Fig. 3A).

Because pneumococcal strains were cultured in THB before being resuspended in the nutritionally poorer CTM for DNA degradation.
degradation, there is a possibility that the impaired DNA degradation in CTM was due to the inability of pneumococcus to adjust to poorer growth conditions. To rule out this possibility, pneumococcal strains were also cultured in CTM to achieve 10^6 CFU/ml, washed and resuspended in THB or CTM medium and examined for DNA degradation. Again, D39, R6, TIGR4, 0100993 and their competence-deficient \( \Delta \text{comD} \) derivatives degraded DNA efficiently in THB, but not in CTM (Fig. 4). These results suggest that EndA-mediated degradation of extracellular DNA is partially dependent on the nutritional condition of growth medium. Collectively, the aforementioned results suggest that competence-independent nucleolytic activity of EndA is conserved in different pneumococcal strains. Furthermore, EndA degrades DNA with equal efficiency in the presence or absence of capsule. However, a richer medium such as THB allows better bacterial growth and higher nucleolytic activities.

Competence-dependent EndA Activity Mediates Rapid Release of Small DNA Fragments Right After CSP1 Stimulation

Contrary to previous reports [7,8], our results indicate that DNA degradation by EndA is not dependent on the development of pneumococcal competence. To determine if the discrepancy is caused by different methods of measuring nuclease activity, we examined DNA degradation mediated by competent and non-competent pneumococcal cells by measuring the EndA-mediated release of small DNA fragments and nucleotides, as previously published [7,8]. Degradation of P-32-labeled hot DNA was compared among \( \Delta \text{comA} \) cells with or without CSP1 stimulation, \( \Delta \text{endA} \) cells or THB control. As expected, substantial release of small DNA fragments and nucleotides was only observed in the \( \Delta \text{comA} \) and \( \Delta \text{comA}+\text{CSP1} \) groups (Fig. 5A–B), indicating that EndA was responsible for DNA degradation. However, the initial rate of DNA degradation differed significantly. After 30 min of incubation, the amount of small DNA fragments and nucleotides released by CSP1-stimulated competent \( \Delta \text{comA} \) cells was 3.9 fold higher than untreated \( \Delta \text{comA} \), as determined by densitometry analysis. In contrast, the amount of small DNA fragments and nucleotides released by the \( \Delta \text{comA} \) group at this time point was only slightly higher than THB group (Fig. 5A–B), similar to previous reports [7,8]. However, by 90 and 150 min post CSP1 exposure, the amount of small DNA fragments and nucleotides released by the \( \Delta \text{comA}+\text{CSP1} \) became indistinguishable than the \( \Delta \text{comA} \) (Fig. 5A–B).

Because densitometry method only measures DNA degradation in a semi-quantitative manner, we repeated the experiments in Figure 3 using pneumococcal cells grown in THB and in CTM, and quantified the release of small DNA fragments and nucleotides by using a scintillation counter. As shown in Figure 6, after 30 min of incubation, nucleotides/small DNA fragments released by \( \Delta \text{comA} \) grown in the CTM supplemented with CSP1 (21 cpm), is much higher than \( \Delta \text{comA} \) grown in the CTM without CSP1 (0.73 cpm). Similarly, at this time interval, nucleotides/small DNA fragments released by \( \Delta \text{comA} \) grown in the THB with CSP1 (41.23 cpm), is significantly higher than \( \Delta \text{comA} \) grown in the THB without CSP1 (20.2 cpm). This indicates that the competence-induced DNA degradation is apparent at 30 min after CSP1 stimulation. After 90, 150 and 210 min of incubation, competence induced DNA degradation is no longer obvious. At these time intervals, nucleotides/small DNA fragments released by \( \Delta \text{comA} \) grown in THB alone increase dramatically, and is statistically indistinguishable from \( \Delta \text{comA} \) grown in THB supplemented with CSP1. In contrast, after 90, 150 and 210 min of incubation, small DNA fragments/nucleotide released by \( \Delta \text{comA} \) grown in CTM was very low (Figure 6). These observations suggest that competence independent nuclease activity is low when \( \Delta \text{comA} \) was cultured in CTM. Collectively, these results suggest the existence of both competence-dependent and competence-independent EndA nucleolytic activities. Competence-dependent EndA mediates rapid release of small DNA fragments right after CSP1 stimulation while competence-independent EndA activity mediates gradual release of small DNA fragments after the peak of competence for genetic transformation. Overall, the competence-dependent EndA activity is relatively weak and transient, and its contribution to DNA degradation is negligible under our experimental conditions.

Secreted form of EndA Contributes Substantially to the “Competence-independent” Activity of EndA

It has been purported that membrane-localized EndA, presumably recruited by the pseudopilus and other competence apparatus, can only gain access and degrade DNA when pneumococcal cells enter competent state (8). However, our experimental data suggest that the competence-independent activity of EndA is responsible for majority of DNA degradation. We hypothesized that pneumococcal cells secrete EndA into the culture medium during growth and contribute to the competence-independent degradation of the extracellular DNA. A small portion of the EndA molecules may have been recruited by the pseudopilus when pneumococcal cells are competent for genetic transformation. To determine whether EndA is secreted, cell-free supernatants of D39 and its isogenic, \( \Delta \text{comD} \) and \( \Delta \text{allyt} \) mutants were collected overtime and assessed for their ability to degrade DNA. As shown in Fig. 7A, the nuclease activities increase in a time-dependent manner in the cell-free supernatants of D39, \( \Delta \text{comD} \) and \( \Delta \text{allyt} \) grown in the THB. In addition, cell-free supernatants from \( \Delta \text{comD} \) and \( \Delta \text{allyt} \) mutants degraded DNA to the same extent as D39, suggesting that competence development and cell lysis do not contribute significantly to the accumulation of secreted EndA.

To further confirm EndA is secreted into the culture medium, bacterial cells and cell-free supernatant were collected from D39 and its isogenic mutants \( \Delta \text{comD}, \Delta \text{allyt}, \) and \( \Delta \text{endA} \) and subjected to SDS-PAGE in a acrylamide gel incorporated with 15 μg/ml of salmon sperm DNA [5]. The gel was then washed with pure water to remove SDS and allow proteins to renature, and examined for DNA degradation after staining with ethidium bromide. As shown in Fig. 7B, EndA activities were detected in both bacterial lysate as well as cell-free supernatants from D39, \( \Delta \text{comD} \) and \( \Delta \text{allyt} \). In contrast, no nucleolytic band was visible in the \( \Delta \text{endA} \). These results strongly suggest that EndA is secreted in a process independent of competence development or partial cell lysis.
EndA Secretion is Conserved in Different Pneumococcal Strains and is Culture Medium Dependent

Next, we compared the EndA secretion by various pneumococcal strains cultured in THB versus in CTM. D39, TIGR4 and 0100993 cultured in THB secreted much higher amounts of EndA when compared to these strains cultured in CTM (Figure 8A). To quantitatively compare the amount of EndA secretion by pneumococcus growing in THB versus in CTM, 1 ml of cell-free supernatant from D39 grown in THB were diluted 10 folds, 100 folds and 1000 folds. Nuclease activities of different diluents were compared against the nuclease activity of undiluted cell-free supernatant from D39 grown in CTM. Each supernatant samples were exposed to hot PCR product and incubated at 37°C for 1 hr. The release of nucleotides/small DNA fragments was quantified by a scintillation counter. As shown in Fig. 8B, cell-free supernatant of D39 cultured in THB produced a nuclease activity of 348 cpm, decreasing to 40.6 cpm, 3.0 cpm and 0.21 cpm in 10 fold, 100 fold and 1000-fold THB diluents, respectively. In contrast, the nuclease activity of undiluted cell-free supernatant of D39 cultured in CTM is 4.2 cpm, which is approximately the 100-fold diluent of the THB supernatant. These results suggest that D39 cells growing in THB release about 100 times more EndA than D39 cells growing in CTM. These observations could partially explain weak nuclease activities when pneumococcal strains were cultured in CTM (Figs. 3–4).

Degradation of NETs by EndA is Independent of Competence and is Inhibited by ATA

EndA has been previously shown to degrade NETs, a host defense mechanism elaborated by neutrophils [11]. We examined whether the EndA-mediated degradation of NETs in pneumococcus is dependent on the induction of the competence regulon. As expected, NETs were intact in RPMI culture medium or ΔendA treated neutrophils (Fig. 9A). In contrast, D39, ΔcomCDE and ΔcglABCDEFG readily digested NETs (Fig. 9A). In addition, degradation of NETs by D39 cells was inhibited by 15 μg/ml ATA (Fig. 9A). Collectively, these results suggest that the degradation of NETs by EndA is independent of competence development.

EndA but not CglABCDEFG is Required for Lung Infection

The DNA uptake apparatus of pneumococcus encoded by the cglABCDEFG operon was previously reported to be required for DNA degradation during competence [8]. To determine whether competent dependent-EndA-mediated DNA degradation is important for virulence, we compared the bacterial burden of D39, ΔendA and ΔcglABCDEFG using an acute pneumonia model of Pneumococcus (Fig. 9B). Furthermore, inclusion of choline chloride, which completely abolishes the function of all the autolysins including LytA, LytB, LytC and ChpD, neither delays nor attenuates DNA degradation. Collectively, these data refute the argument that rapid DNA degradation is caused by the release of EndA mediated by partial autolysis during pneumococcal growth. Rather, we provide strong evidence that EndA is secreted during normal growth of pneumococcus, which contributes substantially to the competence-independent DNA degradation.

Discussion

According to previous studies, nuclease activity of EndA was only detectable in pneumococcal cells competent for genetic transformation [7,8]. Nuclease activity of EndA was also used as an indicator of competent state [7]. Here, we demonstrate the existence of competence-independent EndA activities that is responsible for the majority of the degradation of extracellular DNA and NETs, and is important for virulence. Several lines of experimental evidence support these conclusions: (i) the overall nuclease activity of competence-deficient mutants is comparable to their parental wild-type D39; (ii) addition of exogenous CSP1 that induces competence development contributes minimally to the overall rate of DNA degradation; (iii) partial cell autolysis, which may release EndA into DNA-rich environments, is not required for the nuclease activity; and (iv) secreted form of EndA contributes substantially to competence-independent nuclease activity of pneumococcus.

The aforementioned observations raise an intriguing question: why is it that in some studies [7,8] the nuclease activity of EndA is only detectable during the competent state while we could detect nuclease activity in both competent and non-competent cells? We attribute part of these discrepancies to different strains used for the experiment. For example, the serotype III clinical isolate 0100993 has weaker nuclease activity than that of D39, R6 or TIGR4. This suggests that different pneumococcal strains express and/or secrete EndA at different levels. Another factor that determines EndA activities is the growth medium. We found that nuclease activity of EndA is severely reduced in the CTM, which is a nutritionally poorer medium when compared to the THB. CTM is a preferred medium used in many genetic transformation studies, likely because it allows the integrity of donor DNA to be preserved for a longer period of time. Based on this, we concluded that previous observation of competence-dependent EndA activity [7,8] does not conflict with our observation of competence-independent EndA activity. Competence-dependent EndA activity is transient but detectable for a short period of time right after CSP1 stimulation. In contrast, competence-independent EndA activity predominates after longer incubation in a richer medium like THB, which favors EndA production.

EndA has been reported to be a transmembrane protein [5]. The thickness of D39 cell wall is estimated to be 23 nm [33]. However, according to the 3D modeling of protein structure using the VMD software program, estimated size of the EndA catalytic domain is approximately 4.5 nm (data not shown). It has been suggested that EndA could only gain access to extracellular DNA through the cglABCDEFG operon encoded pseudopilus uptake apparatus [9]. However, our results show that pneumococcal cells deficient in the Cgl proteins rapidly degrade extracellular DNA. A previous study has attributed the DNA degradation in non-competent pneumococcal cells to the release of EndA molecules into the medium by autolysis or other mechanisms [34]. However, our finding shows that the autolysis deficient ΔlytA mutant degrades DNA as rapidly as the parental wild-type D39. Furthermore, inclusion of choline chloride, which completely abolishes the function of all the autolysins including LytA, LytB, LytC and ChpD, neither delays nor attenuates DNA degradation. Collectively, these data refute the argument that rapid DNA degradation is caused by the release of EndA mediated by partial autolysis during pneumococcal growth. Rather, we provide strong evidence that EndA is secreted during normal growth of pneumococcus, which contributes substantially to the competence-independent DNA degradation.

In conclusion, we have shown that the main nuclease activity of pneumococcal EndA is independent of the competence development in THB. Competence-independent activity of EndA is responsible for rapid degradation of extracellular DNA, NETs, and is required for the virulence of S. pneumoniae during lung infection. Therefore, drugs that inhibit EndA, including ATA, could potentially be used to attenuate pneumococcal-mediated degradation of NETs and spread of infection, and reduce horizontal gene transfer. Also, because EndA is secreted and ...
accessible by antibodies, it may serve as an attractive vaccine target.

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Author Contributions

Conceived and designed the experiments: LZ ZZK BAW GWL. Performed the experiments: LZ ZZK GWL. Analyzed the data: LZ ZZK BAW GWL. Contributed reagents/materials/analysis tools: LZ ZZK BAW GWL. Wrote the paper: LZ GWL.

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