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Kinetics and Phospholipid Specificity of Apolipoprotein N-Acyltransferase

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The enzyme apolipoprotein N-acetyltransferase (Lnt) is an integral membrane protein that catalyzes the last step in the post-translational modification of bacterial lipoproteins. Lnt undergoes covalent modification in the presence of phospholipids resulting in a thioester acyl-envelope intermediate. It then transfers the acyl chain to the α-amino group of the N-terminal diacylglycerol-modified cysteine of apoprotein, leading to the formation of mature triacylated lipoprotein. To gain insight into the catalytic mechanism of this two-step reaction, we overproduced and purified the enzyme of Escherichia coli and studied its N-acetyltransferase activity using a novel in vitro assay. The purified enzyme was fully active, as judged by its ability to form a stable thioester acyl-envelope intermediate and N-acylate the apo-form of the murein lipoprotein Lpp in vitro. Incorporation of [3H]palmitate and mass spectrometry analysis demonstrated that Lnt recognized the synthetic diacylglycerol-modified lipopeptide FSL-1 as a substrate in a mixed micelle assay. Kinetics of Lnt using phosphatidylethanolamine as an acyl donor and FSL-1 as a substrate were consistent with a ping-pong type mechanism, demonstrating slow acyl-envelope intermediate formation and rapid N-acyl transfer to the apolipoprotein in vitro. In contrast to earlier in vitro observations, the N-acetyltransferase activity was strongly affected by the phospholipid headgroup and acyl chain composition.

The post-translational modification of proteins by covalent lipid attachment is universal to all organisms. In bacteria, 1–3% of the genome encodes lipoproteins that are anchored to the cell envelope, where they fulfill various key functions, such as maintenance of the cell wall integrity, nutrient uptake, secretion, extra-cytoplasmic protein folding, and virulence (1–7). Lipoproteins are synthesized as prelipoproteins that are translocated across the cytoplasmic membrane via the Sec or the Twin arginine translocation (Tat) machineries (8–10). With the sequential action of phosphatidylglycerol:prolipid transferases and phosphatidylethanolamine-derived thioether-linked diacylglycerol residue, and of prolipoprotein signal peptidase (Lsp), which cleaves the hydrophobic signal peptide to liberate the α-amino group of Cys1. In a final step, apolipoprotein N-acetyltransferase (Lnt) transfers a preferential phosphatidylethanolamine-derived sn-1-acyl chain to the free α-amino group of Cys1, resulting in mature tri-acylated lipoprotein (Fig. 1) (12–15). All three enzymes are essential for growth and viability of Gram-negative bacteria and are located in the inner (cytoplasmic) membrane (16–18). In proteobacteria, N-acylation of lipoproteins is a prerequisite for transfer of lipoproteins to the outer membrane via the Lol machinery (17, 19, 20). The gene encoding Lnt was originally identified in Salmonella enterica sv. Typhimurium and is conserved in proteobacteria and in mycobacteria (21, 22). Recent findings demonstrated that N-acylated lipoproteins exist in various staphylococcal species, suggesting that apolipoprotein N-acetyltransferase activity is present in firmicutes, although an int gene could not be identified in the genomes of this class of bacteria (23, 24).

Initial studies of Escherichia coli Lnt assigned its first enzymatic properties, e.g. a pH optimum of 6.5–7.4, thermostability up to 60 °C, and activity dependence upon the presence of detergents (14). More recently, it was demonstrated that Lnt is anchored in the inner membrane by at least six transmembrane helices with a nitrilase-like periplasmic domain containing the catalytic triad Glu387–Lys333–Cys387 (17, 22). The formation of an extra-cytoplasmic thioester acyl-envelope intermediate suggested that Lnt functions by an initial nucleophilic attack of the activated thiol of Cys387 on the carbonyl group of the sn-1-acyl chain of a phospholipid like phosphatidylethanolamine (PE)3 (25). The by-product of this first step, a lysophospholipid (Fig. 1), is flipped across the inner membrane by the lysophospholipid transporter LplT and is recycled by the bifunctional enzyme 2-acyl-PE acyltransferase/acyl-ACP synthetase (26, 27). In the second step of the Lnt reaction, the acyl-envelope intermediate is resolved by the α-amino group of an incoming apolipoprotein. Mycobacterial lipoproteins are modified with a diacylglycerol moiety containing mycobacterium-specific fatty acids (21). The observation that Lnt of Mycobacterium smeg-

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2 To whom correspondence should be addressed: Molecular Genetics Unit, Institut Pasteur, 25 Rue du Docteur Roux, 75724 Paris Cedex 15, France. Tel.: 33-1-40.61.36.83; Fax: 33-1-45.68.89.60; E-mail: niebud@pasteur.fr.

3 The abbreviations used are: PE, phosphatidylethanolamine; PA, phosphatidic acid; PC, phosphatidylcholine; PG, phosphatidylglycerol; lysophosphatidic acid; DDM, dodecyl β-maltoside; Mal-PEG, maleimide-polyethylenglycol; Tricine, N-[2-hydroxy-1,1-bis(hydroxymethyl)ethyl]glycine; CV, column volume.
**Characterization of Apolipoprotein N-Acyltransferase**

**FIGURE 1. Lipoprotein modification pathway in E. coli.** An unmodified prolipoprotein with a hydrophobic signal peptide (SP) containing the conserved lipobox sequence is first modified by the enzymes phosphatidylglycerol:prolipoprotein diacylglycerol transferase (Lgt) and prolipoprotein signal peptidase (Lsp). This apolipoprotein is further modified by apolipoprotein N-acetyltransferase (Lnt), which attacks the sn-1 acyl chain of a phospholipid, forms a thioester acyl-enzyme intermediate, and transfers the acyl group to the α-amino group of the diacylglycerol-cysteine, leading to a mature triacylated lipoprotein. The acyl donor indicated is based on one of the major E. coli phospholipids, 1-palmitoyl-2-vaccenoyl-sn-glycero-3-phosphoethanolamine (16:0, cis-11,12-18:1).

**TABLE 1**

**Strains and plasmids used in the study**

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<thead>
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<td><em>E. coli</em> K-12 Δ(lac-pro) F’ (lacIΔZ15 pro A15 Tn10)</td>
<td>Laboratory collection</td>
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<tr>
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<td>BW25113 ybeX- (kan-rpoCter-pRNA-lnt)</td>
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<td>plasmids</td>
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*matis* is nonfunctional in *E. coli* suggests that the diacylglycerol group also plays a role in substrate specificity. It is currently not known how Lnt binds its substrates or how it removes the acyl group from phospholipids and subsequently attaches it onto a lipoprotein. We developed an apolipoprotein N-acetyltransferase *in vitro* assay based on purified enzyme, a synthetic lipopeptide substrate and various phospholipid acyl donors. We present its kinetic parameters and determined specificity of fatty acid chains and phospholipid headgroups of apolipoprotein N-acetyltransferase.

**EXPERIMENTAL PROCEDURES**

**Chemicals and Materials**—[9,10-3H]Palmitate, 5 mCi (185 MBq), were purchased from PerkinElmer Life Sciences. The cysteine alkylation reagent mPEG-MAL (5 kDa) was obtained from Creative PEG Works, Winston Salem, NC. All phospholipids used in the study were from Avanti Polar Lipids, Alabaster, AL. The lipopeptide FSL-1-Biotin was obtained from EMC Microcollections, Tübingen, Germany. Plasmid pASK-IBA3+ anhydrotetraacycline, Streptactin®, Sepharose, and desthiobiotin were purchased from IBA GmbH, Göttingen, Germany. The protease inhibitor 4-(2-aminoethyl) benzenesulfonyl fluoride hydrochloride was from Interchim SA, Montluçon, France. All molecular biology enzymes were either from New England Biolabs, Ipswich, MA, or from Fermentas GmbH, St. Leon Rot, Germany. All other chemicals, including oligonucleotides, were either obtained from Sigma or from Carl Roth GmbH, Karlsruhe, Germany, and were at least of analytical grade.

**Bacterial Strains and Plasmids**—All strains and plasmids used in the study are listed in Table 1. PAP105 was used as an *E. coli* strain in all experiments unless indicated otherwise. Plasmid pCHAP9510 was generated by cloning a synthetic DNA fragment (Sigma) encoding the myc tag amino acids in-frame with the Strep tag sequence of pASK-IBA3 using NcoI and PstI restriction sites. The *lnt* gene from *E. coli* was PCR-amplified from genomic DNA using the oligonucleotides Lnt_20a_BsaI (5'-GGATAGGGTCTCAAATGGCTTTTGC-3') and Lnt_20b_BsaI (5'-CACGATGCTACGCGGTCAATAGCACGCAG-3') as primers and cloned into the pASK-IBA3+ using BsaAI restriction sites. The Lnt(K335A) and the single cysteine Lnt(Cys387) variant were obtained by PCR using plasmids pCHAP7650 and pCHAP7556 as templates, respectively (17, 22). PCR fragments were cloned either in pASK-IBA3+ using BsaAI restriction sites (Lnt(K335A)) or in pCHAP9510 (Lnt(Cys387)) using EcoRI and Xhol restriction sites.

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Characterization of Apolipoprotein N-Acyltransferase

Growth Conditions for Overproduction of Lnt—Myc-Strep. E. coli PAP105 with plasmids pCHAP9515, pCHAP9519, or pCHAP9531 was grown in 10 ml of LB with 100 μg ml⁻¹ ampicillin from a single colony at 37 °C overnight. This preculture was used to inoculate 1 liter of LB medium containing 0.1% glucose, and bacteria were initially grown at 30 °C under aeration at 180 rpm on a rotary shaker. At an A₆₀₀ of 0.6–0.8, int expression was induced by the addition of anhydrotriacycline to 100 ng μl⁻¹, and the culture was transferred to 25 °C, which allowed further growth, albeit at a drastically lowered rate. After 4–5 h and at an A₆₀₀ of 1.3–1.6, cells were harvested by centrifugation and washed with buffer P1 (20 mM Tris-HCl, pH 8, containing 150 mM NaCl, and 0.5 mM EDTA). This buffer was used in all subsequent steps, and modifications are as indicated. The pellet was used directly for the preparation of membranes or stored up to 36 h at −25 °C.

Membrane Preparation and Solubilization of Lnt—All steps were carried out at 4 °C. The frozen pellet was resuspended in 25 ml of P1 with 0.1 μg ml⁻¹ DNase and 0.2 mM 4-(2-aminoethyl)benzenesulfonyl fluoride hydrochloride. Cells were broken by three passages through a French pressure cell at 10,000 p.s.i. Unbroken cells and debris were removed by centrifugation at 6,000 × g for 10 min. The membrane fraction was prepared from the supernatant by ultracentrifugation at 120,000 × g for 60 min. The membrane pellets were either stored for up to 48 h at −25 °C or immediately washed and resuspended in 9 ml of P1. Triton X-100 was added to a final concentration of 2% (v/v) to solubilize membrane proteins, and the suspension was incubated for 1 h with intermittent inversion on a rotation apparatus. All insoluble material was removed by centrifugation at 40,000 × g for 30 min.

Protein Purification—Lnt was purified by Strep-Tactin® affinity chromatography using a prepacked Strep-Tactin® Superflow® high capacity column (IBA) coupled to an Äkta Purifier FPLC system (GE Healthcare). The column was equilibrated with 2 column volumes (CV) of P1 containing 2% Triton X-100. After binding 10 ml of the Triton X-100-solubilized membrane fraction, the column was washed with at least 4 CV of P1 containing 0.1% Triton X-100. A second wash with 4 CV of P1 containing 0.05% DDM replaced the Triton X-100 as observed by a decrease in UV absorbance 280 nm. The protein was eluted from the column with 3 CV of P1 containing 0.05% DDM and 2.5 mM desthiobiotin. Purity was judged after SDS-PAGE and staining with Coomassie Brilliant Blue, and the identity of the protein was confirmed by immunoblotting. The protein concentration was determined by measuring the absorbance at 280 nm and by the BCA method using BSA as a standard. Purified protein was stored in the elution buffer at −25 °C for at least 5 months without a detectable loss in activity.

Alkylation with Mal-PEG—The presence of free thiol groups in wild type Lnt and the single cysteine Lnt(C23A,C62A) was determined by their accessibility to the alkylating agent Maleimide-conjugated polyethylene glycol, 5 kDa (Mal-PEG) as described previously (25). Briefly, purified protein was incubated at 37 °C in the presence or absence of phospholipids for 5 min and denatured by adding of 0.5% SDS in 1 M Tris-HCl, pH 8. Mal-PEG was added to a final concentration of 0.2 mM and incubated for 30 min at room temperature. An excess of cysteine (2 mM) was added to terminate the reaction. As the Mal-PEG adds roughly 5 kDa to their molecular mass, alkylated proteins could be identified easily by SDS-PAGE followed by immunoblotting.

Preparation of Apo-Lpp—An int conditional null mutant expressing the int gene from an arabinose-inducible promoter was used to obtain the apo-form of Lpp, essentially as described earlier (17). Lnt was depleted when cells were grown in LB at 37 °C with d-fucose (0.2% (w/v)) to repress the arabinose promoter. After 8 generations (~180 min), growth arrest was observed, and the A₆₀₀ remained constant. Cells were harvested 30 min after the growth arrest, which was accompanied by the accumulation of apolipoprotein. Appearance of the apo-form of Lpp was confirmed by immunoblotting using polyclonal anti-Lpp antibodies. Spheroplasts of these samples were used to isolate the membrane fraction that was either frozen or directly used to analyze Lnt activity.

Lnt in Vitro Assay with Unlabeled Phospholipids—When apo-Lpp was used as a substrate, the assay was carried out using the membrane fraction of Lnt-depleted cells as a source of acyl donor and apolipoprotein substrate. Membranes were added to a final protein concentration of 250 ng μl⁻¹ to the standard reaction buffer composed of 50 mM Tris-HCl, pH 7.2, 150 mM NaCl, and 0.1% Triton X-100. A reaction mixture of 50 μl was briefly sonicated in a sonicator bath and preincubated at 37 °C prior to the addition of purified enzyme at 10 ng μl⁻¹. Samples were removed at time intervals, and Lnt was rapidly inactivated by adding 5% SDS and heating to 100 °C. The relative levels of apo- and mature Lpp were monitored by SDS-PAGE and immunoblotting using anti-Lpp antibodies.

The standard assay for Lnt activity using the lipopeptide FSL-1 was carried out in a reaction volume of 20 μl containing commercial phospholipids and FSL-1 at concentrations of 500 and 5 μM, respectively. Phospholipids were added as mixed micelles in 0.1% Triton X-100. Following a brief sonication and preincubation at 37 °C, 3.5 nM (0.2 ng μl⁻¹) of Lnt was added. To determine the initial rates of the Lnt reaction, samples were taken at least at four time points within the first 120 min of the reaction. The time course of the first (slow) step of the reaction was 0, 5, 10, and 20 min and of the second (fast) step was 0, 2, 5, and 10 min. Samples of 2 μl were removed at time intervals and heat-inactivated in the presence of 5% SDS. Aliquots of the samples were analyzed by SDS-PAGE and immunoblotting.

Preparation of [³H]Palmitate-labeled Membrane Vesicles—Radioactive labeling of whole cells was performed as described previously (23). Cells were grown at 37 °C to A₆₀₀ 0.2, and [³H]palmitate was added to 100 μCi ml⁻¹. After 60 min, cells were harvested by centrifugation and washed with P1 buffer. The total membrane fraction was prepared from spheroplasts as described previously (28). The membranes were heated at 100 °C for 10 min to inactivate endogenous Lnt and stored as 50-μl aliquots at −25 °C.

Lnt Assay with [³H]Palmitate Membrane Vesicles—The assay was performed essentially as described for apo-Lpp, except that [³H]palmitate-labeled membrane vesicles at 200 nCi μl⁻¹ were used as source of phospholipids, and the apolipoprotein substrate was 20 ng μl⁻¹ of FSL-1-Biotin peptide. Following heat
inactivation of Lnt, FSL-1 was purified by its biotin tag using the streptavidin mutein matrix (Roche Applied Science). Purification was carried out according to the manufacturer’s instructions except that 0.1% of Triton X-100 was added to the wash and elution buffers. Elution was performed in three 100-μl fractions that were combined and added to 5 ml of scintillation liquid (GE Healthcare). Radioactivity incorporation into the peptide was determined by measuring \[^3H\]palmitate with the Tri-Carb-2100-TR liquid scintillation analyzer (PerkinElmer Life Sciences) and was expressed as the percentage of total radioactivity in the reaction mixture.

**SDS-PAGE, Immunoblotting, and Quantification**—Purified Lnt was detected after Tris-glycine-SDS-PAGE (10% acrylamide) and immunoblotted using \(\alpha\)-Myc, anti-Strep-tag II, and \(\alpha\)-Lnt antibodies. Rabbit polyclonal antibodies against Lnt were raised using purified His-tagged periplasmic domain of the enzyme as antigen (Genscript). This polypeptide was produced in inclusion bodies in the cytoplasm and was purified by Co\(^{2+}\) affinity chromatography. These results showed that purified Lnt was in its acylated form, detergent-solubilized and delipidated enzyme was treated with Mal-PEG after incubation in the presence or absence of phospholipids. To facilitate analysis of the acylation state of Lnt, a variant in which cysteines at positions 23 and 62 were replaced by alanine was also purified. The variant enzyme Lnt(C23A,C62A) was previously shown to complement fully an int conditional null mutant (25) and to be fully active in vitro (see below). In the absence of phospholipids, Lnt was modified by Mal-PEG (5 kDa), resulting in a slower migrating band in SDS-PAGE (Fig. 2B, lane 2). In the presence of phospholipids, Lnt was protected from alkylation, indicating that the thioester blocked the thiol of the active site cysteine (Fig. 2B, lane 4). Further evidence for the absence of a stable thioester in the purified enzyme was provided by incubating it with \[^3H\]palmitate phospholipids in labeled membrane vesicles. In this assay, Lnt formed a stable thioester acyl-enzyme intermediate that was detected by SDS-PAGE and fluorography (Fig. 2B, lane 4). The identity of the fusion protein was confirmed by immunoblotting using antibodies against the periplasmic domain of Lnt and against the myc and strep tag (Fig. 2B) (data not shown).

The purified enzyme preparations were stored in elution buffer at \(-25^\circ C\) for up to 5 months without significant loss of activity. Overall, this protocol yielded up to 1 mg of protein/liter of cell culture in a single-step purification.

**Purified Lnt Exists Mainly in the Nonacylated Form**—Lnt exists as an extra-cytoplasmic acyl-enzyme intermediate in vivo because of the formation of a stable thioester between the active site cysteine and a palmityl residue, as demonstrated by \[^3H\]palmitate labeling and by cysteine alkylation with Mal-PEG (25). To determine whether purified Lnt was in its acylated form, detergent-solubilized and delipidated enzyme was treated with Mal-PEG after incubation in the presence or absence of phospholipids. To facilitate analysis of the acylation state of Lnt, a variant in which cysteines at positions 23 and 62 were replaced by alanine was also purified. The variant enzyme Lnt(C23A,C62A) was previously shown to complement fully an int conditional null mutant (25) and to be fully active in vitro (see below). In the absence of phospholipids, Lnt was modified by Mal-PEG (5 kDa), resulting in a slower migrating band in SDS-PAGE (Fig. 2B, lane 2). In the presence of phospholipids, Lnt was protected from alkylation, indicating that the thioester blocked the thiol of the active site cysteine (Fig. 2B, lane 4). Further evidence for the absence of a stable thioester in the purified enzyme was provided by incubating it with \[^3H\]palmitate phospholipids in labeled membrane vesicles. In this assay, Lnt formed a stable thioester acyl-enzyme intermediate that was detected by SDS-PAGE and fluorography (Fig. 2C). These results showed that purified Lnt is mainly unacylated but in an active conformation such that it can attack the sn-1-acyl chain of a phospholipid and become acylated.

**Purified Lnt N-Acylates Endogenous Apo-Lpp in Vitro**—Lpp is one of the most abundant proteins in E. coli, exceeding 500,000 molecules per cell (30, 31). Lnt catalyzes the N-acylation of apo-Lpp in vivo, and its correct lipid modification was shown to be essential for cell envelope integrity and for viability (17, 32). Hence, the lipidation intermediates of Lpp have often been used as model substrates to study lipoprotein modification in vitro and in vivo (17, 33, 34). To test whether the detergent-solubilized and -purified Lnt was active under in vitro conditions, we analyzed N-acylation of apo-Lpp in mixed detergent-phospholipid micelles. Apo-Lpp was prepared from a conditional int null mutant (17). In this strain, the int gene is exclu-
sively expressed from an arabinose-inducible promoter. The absence of arabinose leads to depletion of Lnt and accumulation of apo-Lpp (17, 22). The apo-form can be distinguished from the mature (triacylated) form by its faster migration in a high resolution Tricine-SDS-PAGE (32). A mixture of Lpp species corresponding to the mature form, the accumulated apo-form, and Lpp linked to peptides of the peptidoglycan backbone was detected in vesicles prepared from spheroplasts of Lnt-depleted cells (Fig. 3A, 0 min). Upon incubation of Lnt with detergent-solubilized vesicles, apo-Lpp was converted over time into mature Lpp, as indicated by the disappearance of the apo-Lpp (Fig. 3A, 0–45 min). These data indicate that Lnt N-acylated endogenous apolipoprotein *in vitro*.

**FIGURE 2. Purification of Lnt and formation of the acyl-enzyme intermediate *in vitro*.** A, SDS-PAGE of protein fractions at different stages of Lnt purification. Proteins were stained with Coomassie Brilliant Blue and molecular mass standards (M) are given in kDa. Lnt was overproduced and purified from the membrane fraction (Mbr), solubilized in Triton X-100 (TX100), and bound specifically to a Strep-Tactin® column. After washing, purified Lnt was eluted from the column (E). FT indicates flow-through. See “Experimental Procedures” for details. B, alkylation of Lnt by Mal-PEG. Purified Lnt(C23A,C62A) was incubated in the absence (lanes 1 and 2) or presence (lanes 3 and 4) of phospholipids (PL) and subsequently treated with the thiol-specific alkylation reagent Mal-PEG (lanes 2 and 4). Lnt was detected by immunoblotting using α-Myc antibodies. C, purified Lnt forms a palmitoyl enzyme intermediate *in vitro*. Lnt was added to phospholipid vesicles isolated from [3H]palmitate-labeled cells and analyzed by SDS-PAGE and fluorography.

**Lnt N-Acylates a Synthetic Diacylglyceryl Lipopeptide in Vitro**—Initial attempts to study the kinetics of the apolipoprotein N-acyltransferase were hindered by the fact that defined substrates were unavailable and that entire membranes were used as the sources of the enzyme. To overcome this problem, we analyzed whether a synthetic diacylated lipopeptide could be used as a substrate to study Lnt activity. FSL-1-Biotin (FSL-1) is a synthetic peptide based on a lipoprotein from *Mycoplasma pneumoniae* and has often been used as model peptide to study the activation of Toll-like receptors (35, 36). This peptide carries a thioether-linked dipalmitoyl-glyceryl modification on its N-terminal cysteine residue and contains a free α-amino group, resembling perfectly the N terminus of an
apolipoprotein. When FSL-1 was incubated with Lnt and [3H]palmitate-labeled heat-inactivated vesicles in the presence of detergent, more than 10% of total radioactivity was specifically incorporated into FSL-1 (Fig. 3B). The incorporation of [3H]palmitate in FSL-1 was strictly dependent on active Lnt, because [3H]palmitoyl transfer to FSL-1 was not observed when using an inactive Lnt variant in which Lys335 was replaced by Ala (Fig. 3B, dashed line) (22). These data suggested that FSL-1 was N-acylated by Lnt. To confirm that palmitate was attached to FSL-1, an Lnt-dependent reaction containing sn-1-(16:0)-2-(18:1)-PE was analyzed by MALDI-TOF mass spectrometry. The m/z 2246.4 peak corresponding to FSL-1 (average mass accuracy ±30 ppm) was detected both in the absence and in the presence of Lnt. In contrast, the m/z 2484.7 peak was only observed after completion of the reaction with Lnt (Fig. 4). The mass shift of 238.3 between the two peaks demonstrated that Lnt catalyzed the N-palmitoylation of FSL-1 (expected mass of palmitoyl moiety = 238.4 Da). Similar results were obtained by electrospray ionization-MS analyses (data not shown). These data provided direct evidence that the FSL-1 lipopeptide is recognized as an Lnt substrate in vitro and also confirmed that the sn-1-acyl chain (C16:0, 238.4 Da) and not the sn-2-chain (C18:1 and 264.4 Da) is transferred from the phospholipid to the N terminus of apolipoprotein.

Enzymatic Properties of Lnt—The use of FSL-1 and chemically defined phospholipids together with purified Lnt facilitated studies on the substrate and acyl donor specificity and allowed the determination of kinetic parameters. Time-dependent N-acylation using polar lipid extract from E. coli (Fig. 5A) or synthetic phospholipids as acyl donors was analyzed on high resolution Tricine-SDS-PAGE followed by blotting onto nitrocellulose membranes and detection of biotinylated FSL-1 using a streptavidin-peroxidase conjugate. In the absence of enzyme or phospholipids, FSL-1 migrated aberrantly with an apparent size of 5 kDa, despite its theoretical molecular mass of 2246 Da that was confirmed by MALDI and electrospray ionization-MS. Addition of Lnt resulted in a time-dependent appearance of a slightly slower migrating variant of the peptide, corresponding to the N-acylated form of FSL-1 (Fig. 5A). The quan-

![FIGURE 3. Purified Lnt is functional in vitro. A, Lnt N-acylates apo-Lpp in vitro. Purified Lnt was incubated with detergent-solubilized membrane vesicles derived from Lnt-depleted cells containing apo-Lpp as substrate and phospholipids (PL) as acyl donors. Samples were taken at different time points and analyzed by Tricine-SDS-PAGE and immunoblotting using anti-Lpp antibodies. Asterisks indicate Lpp covalently cross-linked to peptidoglycan fragments. B, Lnt-dependent [3H]palmitoyl incorporation in FSL-1. The diacylglycerol-lipopeptide FSL-1 was mixed with phospholipid vesicles isolated from [3H]palmitate-labeled cells. Following the addition of Lnt (○) or inactive Lnt(K335A) (△), the reaction mixture was incubated at 37 °C, and samples were removed at time intervals. The peptide was purified via streptavidin-biotin affinity, and [3H]palmitate incorporation was determined by scintillation counting. The amount of [3H]palmitate incorporated into the peptide is expressed as a percentage of the total amount of radioactivity in the reaction mixture from three independent experiments.

![FIGURE 4. In vitro N-palmitoylation of FSL-1 by Lnt. Comparison of MALDI mass spectra of FSL-1 after the N-acylation reaction in the absence (upper panel) and in the presence (lower panel) of Lnt. The m/z 2200–2600 region of the MALDI spectra is shown, and only monoisotopic significant peaks (S/N ratio >3) are annotated. Y axis represents the relative signal intensity. a peaks correspond most probably to a secondary product of the FSL-1 synthesis (m/z = 2230.38) and its N-palmitoylated form (m/z = 2468.63). b peaks are sodium adducts of FSL-1 and N-palmitoyl-FSL-1 peptides.](http://www.jbc.org/)

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Figure 5. Lnt N-acylation activity with FSL-1. A, Lnt (10 nM) was added to a reaction mixture composed of polar lipid extract from E. coli (500 μM) and FSL-1 (60 μM). Samples were removed at time intervals from the reaction and analyzed by Tricine-SDS-PAGE and immunoblotting against the biotin tag of FSL-1. B, concentration of N-acylated FSL-1 in the reaction is represented as a function of time. C, initial rates of N-acylation activity under standard assay conditions (see “Experimental Procedures”) were strictly dependent on the presence of detergent and are displayed as function of the concentration of Triton X-100.

The existence of an acyl-enzyme intermediate in vivo (Fig. 2B) (25) and in vitro (Fig. 2C) suggested a two-step reaction via a “ping-pong” mechanism, where the initial attack of the phospholipid induces the formation of the intermediate (ping) and is followed by the N-acyl transfer to the apolipoprotein (pong). To verify that kinetics were consistent with this type of mechanism, the acyl donor concentrations were varied with different constant concentrations of FSL-1. When plotting the reciprocal of \( V_0 \) versus the reciprocal of the \( sn-1-(16:0)-2-(18:1) \)-PE concentration, parallel lines were obtained (Fig. 6). These results show that the N-acyl transfer catalyzed by Lnt proceeds via ping-pong mechanism.

Kinetics and Polar Headgroup Specificity—Previous in vivo studies on the site-specific turnover of PE and, more recently, acyl transfer from phospholipids carrying acyl chains selectively labeled with \[^{14}C\]palmitate or \[^{3}H\]palmitate demonstrated that the \( sn-1 \)-acyl chain is the primary acyl donor for Lnt (15, 19). This was further confirmed here by mass spectrometry. The mechanism, however, by which Lnt recognizes phospholipids as acyl donor is still unknown. It has been proposed that Lnt does not show any preference for a particular type of phospholipid with respect to headgroup structure or acyl chain length and composition. This proposal was based on the observation that PE, the major phospholipid in E. coli, was not essential for the N-acylation of lipoproteins in vivo and that all three phospholipids of E. coli (PE, PG, and cardiolipin) could serve as acyl donors in vitro (14, 37). The broad specificity of Lnt was further supported by a nonkinetic analysis using purified Lnt (19).

The in vitro Lnt assay using FSL-1 as an acceptor peptide allowed us to determine N-acylation kinetics in the presence of defined phospholipids. In a first approach, phospholipid-de-
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Accordingly, the $V_{\text{max}}$ value was determined as 350 µmol FSL-1 min$^{-1}$ µmol$^{-1}$ Lnt. Corroborating to a specific activity of 5.6 ± 0.3 µmol FSL-1 min$^{-1}$ mg$^{-1}$ Lnt, PG and PA also served as acyl donors, albeit at reduced rates. Their $K_m$ and $V_{\text{max}}$ values were 10.4 µM and 127 µmol FSL-1 min$^{-1}$ µmol$^{-1}$ Lnt for PG and 3.8 µM and 106 µmol FSL-1 min$^{-1}$ µmol$^{-1}$ Lnt for PA, respectively. The catalytic efficiencies of the enzyme in the presence of these three phospholipids were similar with $k_{\text{cat}}/K_m$ values ranging in the same order of magnitude with 7.3 × 10$^4$ for PE, 4.8 × 10$^5$ for PA, and 2.0 × 10$^5$ for PG. Only trace activity was observed when using PC ($K_m$, 18.2 µM; $V_{\text{max}}$, 27 µmol FSL-1 min$^{-1}$ µmol$^{-1}$ Lnt; and $k_{\text{cat}}/K_m$, 9.2 × 10$^4$ M$^{-1}$ s$^{-1}$). For phosphatidylserine, enzyme activity was below detection limits (data not shown). Variations in the length and composition of the acyl chains on PC did not increase FSL-1 N-acylation. This result provided additional evidence that the polar headgroup plays a crucial role in recognition by Lnt and that differences in micelle-phospholipid structure did not account for reduced activity.

Acyl Chain Specificity—In a second approach, we studied the effect of the acyl chain length and composition on Lnt activity by monitoring the formation of the thioester thio-acyl-enzyme intermediate. After incubation with standard PE carrying sn-1-(16:0) and sn-2-(18:1) acyl chains, Lnt was largely inaccessible to alkylation by Mal-PEG (Fig. 8A, 3rd lane). This result suggested that formation of the acyl-enzyme thioester intermediate was most efficient with this phospholipid, in agreement with optimal enzyme activity observed under these conditions (Figs. 7B and 8B). Similar results were obtained when PE with acyl chains shorter than 16 carbons or with two unsaturated C16 acyl chains were used (Fig. 8A, 4th and 6th lanes). Acyl-enzyme intermediate formation did not occur when sn-1,2-(16:0)-PE, sn-1,2-(18:0)-PE, or branched hydrocarbon chains like archaeal type phytanyl residues (sn-1,2-(16:0;4ME)-PE) were used (Fig. 8A, 5th, 7th, and 8th lanes). To analyze acyl chain-dependent phospholipid recognition by Lnt in more detail, we used the kinetic assay with various concentrations for each type of PE (Fig. 8B). The ability to form a thioester acyl-enzyme intermediate with different types of PE was reflected by their kinetics. Highest activity relative to standard PE (sn-1-(16:0)-sn-2-(18:1), $K_{\text{cat}}/K_m$, 4.8 × 10$^4$ M$^{-1}$ s$^{-1}$; $K_{\text{cat}}$, 462 µM; $k_{\text{cat}}$, 2.2) was observed with PE carrying a saturated C18 on sn-1 and an unsaturated C18 on sn-2 (sn-1-(18:0)-sn-2-(18:1), $k_{\text{cat}}/K_m$, 2.0 × 10$^5$ M$^{-1}$ s$^{-1}$; $K_{\text{cat}}$, 2941 µM; $k_{\text{cat}}$, 6.0). PE with sn-1,2-(12:0) or with sn-1,2-(16:1) acyl chains served as functional acyl donors albeit with reduced efficiencies with $k_{\text{cat}}/K_m$ values of 8.5 × 10$^4$ and 8.9 × 10$^5$ M$^{-1}$ s$^{-1}$, respectively. The corresponding $K_m$ and $k_{\text{cat}}$ values were determined as 476 µM and 0.4 s$^{-1}$ (12:0) and 363 µM and 0.3 s$^{-1}$ (16:1), respectively. Furthermore, sn-1-(16:0) lysophosphatidic acid, which lacks an sn-2 acyl chain, did not serve as a donor. As expected, activity with sn-1,2-(16:0)-PE, sn-1,2-(18:0)-PE, and sn-1,2-(16:0;4ME)-PE was below detection limits (data not shown). These data indicate that also the nontransferred sn-2 acyl chain plays an important role in acyl donor recognition.

DISCUSSION

The enzyme apolipoprotein N-acyltransferase (Lnt) catalyzes the phospholipid dependent N-acylation of bacterial lipo-
proteins, resulting in mature triacylated protein (14, 19). To gain deeper insight into this unique two-step reaction mechanism, we developed a quantitative in vitro assay that allowed us to determine kinetic parameters and acyl donor specificity. In cell extracts, Lnt exists mainly as a thioester acyl-enzyme intermediate with a stable thioester link between a fatty acid residue and the active site cysteine (22, 25). The purified enzyme was mainly nonacylated, as indicated by the fact that the active site thiol was accessible to alkylation by Mal-PEG. It is likely that the acyl-enzyme intermediate resolves during purification as a result of the lipid-detergent exchange. The purified enzyme could reform a stable thioester in the presence of phospholipids as demonstrated by [3H]palmitate labeling and cysteine inaccessibility to alkylation. Auto-acylation of Lnt and the formation of mature Lpp in vitro illustrated that Lnt remained in an overall active conformation upon purification and reconstitution in detergent micelles. We used the small dipalmitoylglyceryl-modified lipopeptide FSL-1 as a substrate to study the kinetics of N-acylation. The determined V_max value for Lnt of E. coli led us to predict that 100–200 molecules of enzyme would be sufficient to achieve the N-acylation of all Lpp produced per cell (500,000) during one generation of growth suggesting that the enzyme is of low abundance but is highly efficient. Only few described membrane-bound acyltransferases use phospholipids as acyl donors. The lecithin-retinol acyltransferase transfers the sn-1-acyl chain from PC to retinol and, like Lnt, proceeds through an acyl-enzyme intermediate (40, 41). PC-dependent N-acyltransferase activity was reported for a very similar enzyme, the lecithin-retinol acyltransferase-like protein RLP-1, which is involved in the synthesis of precursors for bioactive N-acylethanolamines (42). The specific activities reported for both enzymes are at least 1 or 2 orders of magnitude lower than those determined for Lnt (42, 43). A prominent example of another membrane-bound bacterial acyltransferase is PagP. This enzyme is an integral outer membrane protein that transfers the sn-1-palmitoyl chain derived from PE to a lipid A molecule with 2 units of 3-deoxy-d-manno-2-octulosonic acid linked at
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the 6’ position (44). When sn-1-(16:0)-sn-2-(18:1)-PE was used as an acyl donor in the presence of Triton X-100, PagP catalyzed the O-acylation of lipid A at a rate of 7.22 \( \text{mmol min}^{-1} \text{mg}^{-1} \text{PagP} \) (45). Using the same phospholipid as an acyl donor, similar kinetics (5.6 \( \text{mmol min}^{-1} \text{mg}^{-1} \text{Lnt} \)) were determined for the N-acyltransferase activity of Lnt. Despite a completely different second step reaction, amide versus acyloxyacyl formation, both acyltransferases showed comparable enzymatic activities that are well reflected by the high abundance of their substrates, 5 \( \times \) 10^6 for Lpp (30, 31) and 2 \( \times \) 10^6 for lipid A (46).

Lnt was reported to accept a wide range of phospholipid acyl donors in vivo (15, 37). Here, we addressed its acyl donor specificity in more detail using the in vitro assay. PE was most efficient of the phospholipids tested in the N-acyltransferase reaction, which is not very surprising, because PE was reported to be the preferred acyl donor of Lnt in vivo (15). In exponentially growing E. coli cells, PE constitutes between 70 and 80% of cellular phospholipids, making it by far the most abundant component of the membrane (38). Both PG and PA supported apolipoprotein N-acyltransferase activity, although less efficiently than PE. PG is the second most abundant E. coli phospholipid (\(~20\%\)), and a mutant lacking PE apparently N-acylated lipoproteins normally, providing evidence that PG can substitute for PE in vivo (37). PA, a minor component of E. coli membranes (\(~0.2\%)\), is a key intermediate in phospholipid metabolism that accumulates in the absence of PE and is thought to compensate for it in the recruitment of peripheral membrane proteins (49). Phospholipids carrying small headgroups (PE and PG) are preferred over those with more bulky ones (phosphatidylserine and PC). Phospholipid recognition by Lnt was also dependent on the length and composition of the acyl chains. Enzyme activity and acyl enzyme formation was significant and independent of the length and composition of the acyl chains.

In vivo

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