Human 60-kDa Lysophospholipase Contains an N-terminal L-Asparaginase Domain That Is Allosterically Regulated by L-Asparagine

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Background: The 60-kDa human lysophospholipase comprises an N-terminal domain with predicted, yet uncharacterized 1-asparaginase activity and a C-terminal ankyrin repeat-like domain.

Results: The N-terminal domain, termed hASNase1, was identified as a functional structural unit possessing catalytic activity.

Conclusion: hASNase1 is an allosterically regulated bacterial-type cytoplasmic L-asparaginase.

Significance: Domains of multifunctional human proteins harbor homologs of prokaryotic enzymes displaying similar structural and kinetic features.

The structural and functional characterization of human enzymes that are of potential medical and therapeutic interest is of prime significance for translational research. One of the most notable examples of a therapeutic enzyme is L-asparaginase, which has been established as an antileukemic protein drug for more than four decades. Up until now, only bacterial enzymes have been used in therapy despite a plethora of undesired side effects mainly attributed to the bacterial origins of these enzymes. Therefore, the replacement of the currently approved bacterial drugs by human homologs aiming at the elimination of adverse effects is of great importance. Recently, we structurally and biochemically characterized the enzyme human L-asparaginase 3 (hASNase3), which possesses L-asparaginase activity and belongs to the N-terminal nucleophile superfamily of enzymes. Inspired by the necessity for the development of a protein drug of human origin, in the present study, we focused on the characterization of another human L-asparaginase, termed hASNase1. This bacterial-type cytoplasmic L-asparaginase resides in the N-terminal subdomain of an overall 573-residue protein previously reported to function as a lysophospholipase. Our kinetic, mutagenesis, structural modeling, and fluorescence labeling data highlight allosteric features of hASNase1 that are similar to those of its Escherichia coli homolog, EcASNase1. Differential scanning fluorometry and urea denaturation experiments demonstrate the impact of particular mutations on the structural and functional integrity of the L-asparaginase domain and provide a direct comparison of sites critical for the conformational stability of the human and E. coli enzymes.

L-Asparaginases (EC 3.5.1.1; L-asparagine amidohydrolase; L-ASNase2) are enzymes that primarily catalyze the conversion of L-asparagine (L-Asn) to L-aspartic acid (L-Asp) and ammonia, although some of them are able to also hydrolyze L-glutamine (L-Gln) to L-glutamic acid (L-Glu) and ammonia. These enzymes are present in bacteria to mammals and play pivotal roles in amino acid metabolism (1, 2). Enzymes with L-asparaginase activity can be generally classified into two evolutionary distinct families: the bacterial-type and the plant-type L-asparaginases, which are characterized by different structural and biochemical features (3, 4). The bacterial-type enzymes have been further grouped into type I and type II depending on their cellular localization. Type I includes cytosolic enzymes that exhibit low affinity (millimolar $K_{\text{m}}$) for L-Asn, whereas type II enzymes are localized in the periplasm and show considerably higher affinity (micromolar $K_{\text{m}}$) for L-Asn (5). These enzymes have been studied extensively over the last 40 years mainly because two of the type II isoforms (L-ASNases from Escherichia coli and Erwinia chrysanthemi encoded by the ansB genes) serve as therapeutics for the treatment of acute lymphoblastic leukemia (6–8). Conversely, the less studied plant-type L-asparaginases belong to the so-called N-terminal nucleophile (Ntn) hydrolases, which were defined as a new protein structural family in 1995 (9). A major characteristic of this Ntn hydrolase superfamily is a post-translational processing step that generates the active enzyme. The enzymes are expressed as inactive precursors that undergo a slow intramolecular auto-proteolytic cleavage reaction at a specific site resulting in two tightly associated subunits, $\alpha$ and $\beta$, also called protomers. The catalytic residue acting as the nucleophile is exposed at the very N terminus of the newly generated $\beta$-subunit, which remains complexed with the $\alpha$-subunit during catalysis (10–12). A well characterized mammalian member of the Ntn nucleophile hydrolase superfamily is the human lysosomal aspartylglucosa-

L-asparaginase 1; RnASNase1, R. norvegicus L-asparaginase, 1ScASNase1, S. cerevisiae L-asparaginase 1; ansA, gene name for cytoplasmic bacterial L-asparaginases; ansB, gene name for periplasmic bacterial L-asparaginases; Ntn, N-terminal nucleophile; palmitoyl-lysoPC, 1-palmitoyl-sn-glycero-3-phosphocholine; lysoPL, lysophosphatidylinositol; ADIFAB, acrylodated intestinal fatty acid-binding protein; SUMO, small ubiquitin modifier; Ni-NTA, nickel-nitrilotriacetic acid; CAPSO, 3-(cyclohexylamino)-2-hydroxy-1-propanesulfonyl acid.
Allosterically Regulated Human L-Asparaginase

EXPERIMENTAL PROCEDURES

Materials—Yeast extract, peptone from casein, NaCl, Nessler’s reagent, urea, SYPRO Orange, L-asparagine, L-glutamine, L-aspartic acid, glutamate dehydrogenase, α-ketoglutarate, NADH, synthetic 1-palmityl-sn-glycero-3-phosphocholine (palmitoyl-lysophosphatidylcholine, lysoPC), and glycine (glycine max, highly polyunsaturated soybean oil; mixture of different fatty acids and enriched with stearic and palmitic acids), phospholipase A₂ from bovine pancreas, L-α-phosphatidylcholine from soybean (mixture of different fatty acids), and palmitic acid were purchased from Sigma-Aldrich-Fluka. The acrylated intestinal fatty acid-binding protein (ADIFAB) assay kit was obtained from FFA Sciences (San Diego, CA). Dialysis membranes and Coomassie Brilliant Blue G-250 (Bradford reagent) were from Roth (Karlsruhe, Germany). Slide-A-Lyzer was from Pierce. Oligonucleotides were synthesized by IBA GmbH (Goettingen, Germany). Restriction enzymes and T4 DNA ligase were obtained from New England Biolabs (Ipswich, MA). KAPA HiFi polymerase and all PCR reagents were from PeqLab (Erlangen, Germany). Gel extraction and PCR product purification kits as well as nickel-agarose (Ni-NTA) for protein purification were purchased from Macherey Nagel (Düren, Germany). Plasmid purification kits were from Fermentas (Thermo Fisher Scientific, Germany). Genomic DNA preparation kits were from Qiagen (Hilden, Germany).

Cloning of hASNase1, EcASNase1, and Chaperonin 10—The cDNA region coding for the N-terminal domain (termed hASNase1) of the full-length 60-kDa human lysophospholipase was PCR-amplified using as template a cDNA clone of the ASPG gene (human L-asparaginase homolog, NCBI Reference Sequence NM_001080464; GenBank™ accession number BC035836) isolated from female ovarian tissue (Source Biosciences, UK). Four different C-terminal truncation constructs of hASNase1 were generated and subsequently tested for solubility and enzymatic activity as described in the following sections. Ndel and BamHI sites, respectively, were incorporated in the 5′-ends of the amplifying oligonucleotides (primers are listed in the supplemental table). The PCR mixture in a 50-μl final volume consisted of 50 ng of template plasmid DNA, oligonucleotide mixture (10 pmol each), KAPA high fidelity buffer, dNTPs (0.2 mM each), and 1 unit of KAPA HiFi DNA polymerase. The PCR fragment was gel-purified and subjected to additional overlap extension PCRs aiming at the elimination by silent mutation of an internal BamHI restriction site (nucleotide sequence position 740) whose presence would limit the unique cleavage at the 3′-end. The final PCR product was digested with Ndel and BamHI and ultimately ligated overnight at 16°C into the pET14b-SUMO vector (10) using T4 DNA ligase. The ligation mixture was used to transform DH5α E. coli cells. Plasmid DNA isolated from single colonies was digested with Ndel and BamHI to identify positive clones, some of which were sequence-verified. The final constructs include an N-terminal six-histidine tag (His₆) followed by the small ubiquitin modifier (SUMO; Smt3p) tag, which has been shown to improve heterologous protein solubility and stability (26). For bacterial expression, the E. coli BL21(DE3) pLysS strain was transformed with
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each of the four hASNase1 constructs. Expression vectors for EcASNase1 and Chaperonin 10 (GroES; encoded by the Cpn10 gene) were constructed analogously.

Mutagenesis of hASNase1 and EcASNase1—All hASNase1 single site mutants (T19A, T116A, T187S, T187A E266R, E266S, and C299S) and EcASNase1 mutants (R240E and C273S) were generated by applying the QuickChange methodology (Stratagene) using as templates the cloned wild-type genes and KAPA HiFi DNA polymerase. Mutants T187S and T187A were generated in a similar way (the primers are listed in the supplemental table). Site-specific mutations were confirmed by sequencing the entire coding regions (Seqlab, Goettingen, Germany). All hASNase1 mutants were expressed and purified following the protocol described below for the wild-type enzyme.

Protein Expression and Purification of hASNase1, EcASNase1, and GroES—E. coli BL21(DE3) pLysS cells containing the hASNase1 plasmid were cultured overnight at 37 °C in LB medium supplemented with 200 µg/ml ampicillin and 35 µg/ml chloramphenicol. A fraction of this culture (dilution 1:100) was used to inoculate fresh 2× YT medium (1% yeast extract, 1.6% Tryptone, 0.5% NaCl) supplemented with 200 µg/ml ampicillin and 35 µg/ml chloramphenicol. When the bacterial culture reached an A600 of 0.5–0.7, its temperature was lowered to 22 °C, and protein expression was induced by adding isopropyl-1-thio-β-D-galactopyranoside to a final concentration of 0.25 mM. After incubation at 16 °C for 18 h, the cells were centrifuged at 4,000 × g for 30 min, resuspended in Ni-NTA-agarose affinity matrix binding buffer A (50 mM Na2HPO4, 0.3 mM NaCl, 10 mM imidazole, 2% glycerol, 1 mM PMSF, pH 8.0), and ultimately lysed by sonication. The cell lysate was centrifuged at 17,200 × g for 45 min, and to the cleared supernatant was added a fraction (1:10 dilution) of 10× ATP buffer B (50 mM NaH2PO4, 0.3 mM NaCl, 100 mM ATP, 200 mM MgCl2, 500 mM KCl, pH 8.0). The combined mixture was incubated in a 37 °C water bath for 15 min, mixed with pre-equilibrated Ni-NTA-agarose beads, and incubated at 4 °C for 3 h under rotation. Subsequently, the mixture was transferred to a 12-ml polystyrene column and dried by gravity. To remove unspecifically adsorbed material, the nickel resin was first washed with 25 bed volumes of washing buffer C (50 mM Na2HPO4, 0.5 mM NaCl, 20 mM imidazole, 2% glycerol, pH 8.0). The resin was then mixed with a 10-fold diluted buffer B supplemented with GroES at a final concentration of 50 µM. The mixture was incubated at 4 °C for 2 h under rotation, dried by gravity, and then further washed with 10 bed volumes of buffer B without GroES. Finally, the bound His6-SUMO-hASNase1 protein was eluted from the column by applying 300 mM imidazole in buffer A, and fractions were collected dropwise and tested for l-asparaginase activity. All purification steps were performed at 4 °C. The fractions that contained active hASNase1 were pooled, supplemented with GroES at a final concentration of 50 µM and tested for l-asparaginase activity.

For steady-state kinetic analysis, l-Asn concentrations in the range of 0–10 Km were tested in a final volume of 1 ml of 50 mM Tris-Cl, 100 mM NaCl, pH 8 at 37 °C. The final enzyme concentration was ~0.5 µM (20 µg in 1 ml). The obtained V/E (velocity/total enzyme concentration) values were plotted against the respective substrate concentrations. Kinetic constants S0.5 and kcat were calculated from the resulting plots by non-linear regression using the Hill equation (Equation 1) and analyzed by the SoftZymics software (Igor Pro, Wavemetrics).

\[
V = \frac{V_{max}[S]^n}{[S]^{0.5} + [S]^n}
\]  

(Eq. 1)

Here, , V is the initial velocity of the enzymatic reaction, Vmax is the maximal velocity, [S] is the substrate concentration, [S0.5] is
the substrate concentration that yields half-maximal velocity, and \( n \) is the Hill coefficient.

Potential lysophospholipase activity of hASNase1 was tested using a continuous fluorescence assay as described previously (31). Briefly, this assay detects free fatty acid molecules, which are released upon hydrolysis of lysophospholipids by lysophospholipases; free fatty acid then is complexed with the intestinal fatty acid-binding protein conjugated with the fluorescent probe acrylodan, commercially known as a free fatty acid indicator and abbreviated as ADIFAB. Free fatty acid binding to ADIFAB induces a fluorescence signal decrease due to changes of the position of the acrylodan fluorophore, which is located close to the free fatty acid binding pocket of intestinal fatty acid-binding protein. In the present study, we tested palmitoyl-lysoPC and lysoPI as candidate substrates of hASNase1. The assay was performed in a quartz cuvette according to the manufacturer’s instructions using final concentrations of 100 \( \mu M \) for each substrate, 1 \( \mu M \) ADIFAB, and 1.8 \( \mu M \) enzyme (15 \( \mu g \)) in 200 \( \mu l \) of 50 \( mM \) HEPES, 140 \( mM \) NaCl, 5 \( mM \) KCl, 1 \( mM \) \( Na_2HPO_4 \), pH 7.5. Fluorescence intensity was recorded continuously for 10 min at 25 \( ^\circ C \) using a Jasco FP 8300 spectrofluorometer in the high sensitivity mode (excitation at 386 nm and emission at 432 nm with 2.5-nm bandwidths). The assay was initially standardized by testing the fluorescence signal decrease upon binding of palmitic acid (one of the expected products of substrate hydrolysis; lysoPI contains a mixture of fatty acid derivatives; see “Materials”) with the ADIFAB sensor protein as well as by measuring the activity of phospholipase \( A_2 \) against \( \alpha \)-phosphatidylcholine (also provided as a mixture of different fatty acid derivatives). Both hASNase1 and EcASNase1 were tested for lysophospholipase activity.

**Homology Modeling of hASNase1**—A homology model of the N-terminal \( \alpha \)-asparaginase domain of the 60-kDa human lysophospholipase based on the recently determined structure of the homologous EcASNase1 enzyme (25) was obtained from the protein structure prediction service Phyre (32) (Job code 77beadcf4ff84dfc, Fold Library ID c2p2dA, identity of 47%, estimated precision of 100%). The resulting hASNase1 structure was overlaid on the EcASNase1 crystal structure (Protein Data Bank code 2HIM) using PyMOL (33). The putative allosteric and catalytic sites of hASNase1 were analyzed for the presence of critical residues previously identified in EcASNase1.

**Probing the Putative Allosteric Site of hASNase1 by Fluorescence Labeling of Cys\(^{299}\)**—Cysteine residue Cys\(^{299}\) of the catalytically inactive T19A mutant of hASNase1 was labeled using the compound Atto 465-maleimide as a fluorescent dye (Sigma-Aldrich). Maleimides show excellent reactivity with the compound Atto 465-maleimide as a fluorescent dye (34) that has been shown (34) that labeled proteins migrate more slowly than unlabeled species.

For measuring interactions of hASNase1 with \( L \)-Asn, \( L \)-Asn in the concentration range of 0.1–40 \( mM \) was incubated with labeled enzyme (1.5 \( \mu M \) final concentration) in a final volume of 100 \( \mu l \). The samples were left for 2 h on ice to equilibrate. Fluorescence intensity was recorded using a Jasco FP 8300 spectrofluorometer in the high sensitivity mode (excitation at 463 nm and emission at 508 nm with 2.5-nm bandwidths). All measurements were done in triplicate. Because hASNase1 contains overall 6 cysteines, the mutant C299S was also labeled and served as a negative control (background fluorescence) for signal changes upon \( L \)-Asn binding. The resulting saturation binding curve was fitted using SoftZynics software (Igor Pro, WaveMetrics) according to Equation 2,

\[
F = F_0 - \frac{F_{\text{max}}S}{K_d + S}
\]  

(\text{Eq. 2})

where \( F \) is the fluorescence intensity, \( F_0 \) is the fluorescence intensity at zero concentration of ligand, \( F_{\text{max}} \) is the plateau fluorescence intensity, \( K_d \) is the dissociation constant, and \( S \) is the ligand concentration.

**Thermodynamic Characterization of hASNase1 Conformational Stability**—The conformational stability of hASNase1 and EcASNase1 was studied by monitoring changes in intrinsic fluorescence of the stepwise urea-denatured proteins (35, 36). The final urea concentrations ranged from 0.5 to 8 \( M \) using a stock solution of 10 \( M \) urea dissolved in 50 \( mM \) Tris-Cl, 100 \( mM \) NaCl, pH 8. The final concentration of the enzymes was adjusted to 50 \( \mu g/ml \) (\( \sim 1.25 \mu M \) for both enzymes) in a final volume of 100 \( \mu l \). The two tryptophan residues of hASNase1 (Trp\(^{93}\) and Trp\(^{150}\)) exhibited intrinsic fluorescence upon excitation at 295 nm (this wavelength was chosen to reduce excitation of tyrosines), showing an emission maximum at 343 nm. The same excitation and emission maxima were obtained for EcASNase1 containing a single tryptophan residue (Trp\(^{68}\)). Before measurements, equilibrium of the unfolding reactions was confirmed by monitoring the fluorescence signals at different time points until no further change was observed. Data were analyzed assuming a two-state model for reversible protein unfolding. Isothermal urea-induced unfolding experiments were carried out at 10, 15, 20, 25, 30, 35, and 40 °C. For each temperature, data analysis using the linear extrapolation method (37) yielded three parameters: (i) the difference of Gibbs free energy between the native and the unfolded state of the protein (\( \Delta G_{\text{H}_2\text{O}} \)); (ii) the dependence of \( \Delta G_{\text{H}_2\text{O}} \) on denaturant concentration (parameter \( m \), which is the slope of the equation \( \Delta G = \Delta G_{\text{H}_2\text{O}} - m\text{[urea]} \)), which can be thought of as a measure for the sensitivity of the protein toward the unfolding agent; and (iii) the concentration of urea at which the protein is half-unfolded (\( C_{1/2} \)). Subsequently, the calculated \( \Delta G_{\text{H}_2\text{O}} \) values (free energy change at zero concentration of urea) resulting from each temperature were fitted to the Gibbs-Helmholtz equation (Equation 3) (37) as a function of temperature to obtain the \( \Delta C_p \) values. \( T_m \) and \( \Delta H_m \) values as fitting parameters were
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calculated from temperature-induced denaturation experiments (range, 15–85 °C) following the intrinsic tryptophan fluorescence changes of protein similar to urea experiments.

\[
\Delta G(T) = \Delta H_m(1 - T/T_m) - \Delta C_p(T_m - T) + T \ln(T/T_m)
\]

(Eq. 3)

\(\Delta G(T)\) is the Gibbs free energy difference between the folded and unfolded state at temperature \(T, T_m\) is the melting temperature, \(\Delta H_m\) is the enthalpy difference at \(T_m\), and \(\Delta C_p\) the heat capacity difference of the folded and unfolded states at constant pressure.

*Thermal Stability Determination of Wild-type and Mutant hASNase1 Enzymes by Differential Scanning Fluorometry*—The effect of point mutations on thermal stability of hASNase1 was studied by applying differential scanning fluorometry techniques (38, 39). Enzyme samples were dialyzed (Slide-A-Lyzer, 10,000 molecular weight cutoff) against 50 mM Tris-Cl, 0.1 NaCl, pH 8 to remove glycerol and subsequently mixed with SYPRO Orange (Sigma-Aldrich) in a final volume of 20 μl. The final concentrations of the enzyme and the dye were 2 μM and 10% (v/v), respectively; the DMSO stock solution of the dye (5,000×) was prediluted in H2O, giving a 100× solution, from which aliquots were used according to the experimental needs. The samples were mixed in a 96-well plate suitable for real time PCR measurements, centrifuged at 500 rpm for 30 s, and finally sealed with heat-resistant membranes (Microseal B adhesive sealer) to prevent evaporation. The enzyme melting experiments were performed using a CFX96 real time PCR machine (Bio-Rad) with the following settings: 2-min prewarming step at 30 °C and subsequent gradient between 31 and 95 °C with 1 °C/min increments. SYPRO Orange fluorescence was monitored using the filters provided with the machine: FAM (492 nm) for excitation and ROX (610 nm) for emission. Data were exported as an Excel-based worksheet and further analyzed by Igor Pro (Wavemetrics). Melting temperatures (\(T_m\)) were obtained by plotting the first derivative d(AFU)/d\(T\) (where AFU represents arbitrary fluorescence units) of the raw data as a function of temperature increase (40).

*Effects of pH, Divalent Metal Ions, and DTT on Enzyme Activity and Stability*—Effects of pH on enzymatic activity and stability were assessed in a broad range of pH values using several buffers: sodium acetate (pH 3–5), sodium phosphate (pH 6–7), Tris-Cl (pH 7–8.5), and CAPSO (pH 9–10), all at 50 mM concentration in the presence of 100 mM NaCl. For the investigation of the pH effect on the activity of the enzyme, ~50 μg of enzyme were assayed in a final volume of 1 ml at saturating substrate concentration (20 mM L-Asn) applying the Nesslerization method. The discontinuous method was preferred to the NADH-dependent continuous assay because the different pH values could influence the activity of the auxiliary glutamate dehydrogenase enzyme, resulting in unforeseeable artifacts. The stability of hASNase1 under various pH conditions was studied by incubating the enzyme (0.1 mg/ml final concentration) in several buffers for 12 h at 4 °C. Aliquots of these mixtures were assayed for residual L-asparaginase activity by Nesslerization using as standard buffer the one at which the enzyme showed the highest activity based on the previous activity analysis. To study metal ion effects, purified hASNase1 (0.1 mg/ml) was preincubated with a final concentration of 1 mM for each of several divalent metal ions, EDTA, and DTT for 3 h at 4 °C. Subsequently, aliquots were tested for residual activity by the Nesslerization method.

**RESULTS**

*Expression and Purification of the hASNase1 Domain*—Four different C-terminal truncations (numbered 1, 2, 3, and 4; see Fig. 2A) of full-length human lysophospholipase (gene code ASPG; asparaginase homolog) were designed based on sequence alignments of the N-terminal putative L-asparaginase domain and different procaryotic (E. coli, EcASNase1, and Pyrococcus horikoshii, PhASNase1) and eucaryotic (Rattus norvegicus, RnASNase1, and Saccharomyces cerevisiae, ScASNase1) cytoplasmic L-asparaginases (Fig. 1). All four constructs were tested for protein expression in E. coli and solubility under a spectrum of different conditions (culture medium, E. coli strain, isopropyl 1-thio-β-n-galactopyranoside concentration, and temperature). Interestingly, truncation 3, which appears to fit best to the bacterial cytoplasmic enzymes according to the amino acid sequence comparison, was not soluble, and trials to purify the enzyme under denaturation/renaturation conditions (unfolding by up to 8 M urea and refolding by dialysis) ultimately resulted in a catalytically inactive protein. Similarly, truncations 1 and 2 failed to produce soluble, active hASNase1. In contrast, truncation 4 (369 amino acid residues), containing 5 additional amino acids (VEERR) downstream of truncation 3, produced the protein in soluble form (Fig. 2B), and most importantly, the protein purified under native conditions exhibited L-asparaginase activity.

However, when the protein purity was analyzed by SDS-PAGE after the last gel filtration step (see “Experimental Procedures”), a prominent band with an apparent molecular mass of ~60 kDa and an estimated stoichiometric 1:1 ratio to the hASNase1 band was constantly observed (Fig. 3). Multiple attempts to remove this associated protein species by altering standard purification protocols were unsuccessful. This prompted us to analyze this protein band by mass spectrometry by which it was finally identified as the endogenous E. coli chaperone GroEL. Indeed, contamination by co-purified chaperones can be a hurdle in recombinant protein expression and purification setups (41). Although it has been described previously (42) that incubation of the crude cell extract with 10 mM ATP at 37 °C and washing of the affinity column with an ATP-containing buffer facilitate the removal of GroEL, this approach did not prove beneficial in our case. Moreover, attempts to use an engineered E. coli strain (43) (a kind gift from Prof. Ulrich Hartl’s laboratory, Max Planck Institute for Biochemistry, Munich, Germany) carrying the GroEL/ES operon under an arabinose-inducible promoter, which allows tuned regulation of the expression levels of these chaperones, failed to produce the hASNase1 protein (data not shown). Ultimately, we managed to remove the co-purified chaperone by including in the washing buffer the natural binding partner of this protein, GroES, at ~10-fold excess over GroEL (44). It appears reasonable to assume that the presence of ATP-Mg2+-GroES weakened the binding between GroEL and hASNase1, thus facilitat-
ing their separation upon exhaustive washing (Fig. 3). It is
worth mentioning that the bound GroEL did not affect the
activity of hASNase1 because activity measurements on both
enzyme preparations (with and without bound GroEL) yielded
similar kinetic constants.

We emphasize that all hASNase1
variants produced in this work were expressed at levels similar
to the wild-type enzyme and showed no aggregation or precip-
itation tendency at any purification step. GroEL contamination
was present during purification of all mutant enzymes as well.

Size exclusion chromatography experiments allowed us not
only to obtain a highly pure protein as evidenced by SDS-
PAGE analysis (Fig. 3) but also to gain insight into the oligo-
meric state of hASNase1. Strikingly, when we first analyzed
native hASNase1, we observed that the enzyme (\( \approx 40 \text{ kDa} \)) ran
ning as a monomer (molecular mass, \( \approx 40 \text{ kDa} \)) on a Super-
dex 200 gel filtration column (Fig. 4C). As the shift between
oligomeric states of enzymes can be drastically influenced by
their substrates and/or other interacting partners (45, 46), we
decided to perform gel filtration analysis in the presence of the
substrate of hASNase1. Interestingly, with 20 mM L-Asn in the
running buffer, we witnessed a slightly shifted shoulder peak of
the previously observed chromatographic peak assigned to the
40-kDa hASNase1 monomer, indicative of a higher molecular
weight species, although the monomer remained the predomi-
nant form (Fig. 4D). This finding suggested that L-Asn could
trigger the association of monomeric hASNase1 molecules and
consequently induce the formation of dimers and even tetra-
mers, which are the characteristic molecular species reported
for EcASNase1 (25) as we confirmed in the present study (Fig.
4B). However, at enzymatic assay conditions, hASNase1 (\( \approx 0.5 \text{ mM} \)) is expected to be monomeric because this concentration is
6-fold lower than the concentration of the eluted monomeric
protein shown in Fig. 4 (0.5 mg of enzyme loaded on the gel
filtration column eluted in a final volume of \( \approx 4 \) ml of buffer,
\( \text{i.e.} \approx 3 \text{ mM} \) final concentration of the enzyme).

The hASNase1 Enzyme Shows Non-Michaelis-Menten Ki-
netics for L-Asn Hydrolysis Similar to Its E. coli Homolog but
Lacks Lysophospholipase Activity—The N-terminal domain
of human 60-kDa lysophospholipase, which we have designated
as hASNase1, shares 47% sequence identity with
E. coli
cytoplasmic
L-asparaginase (encoded by the
ansA
gene; Ec
ASNase1). The
crystal structure of
Ec
ASNase1 has been reported recently (25),
shedding light on distinct structural features associated with
L-Asn binding to the catalytic site and to the site responsible for
allosteric regulation. Based on this structure, we modeled the
hASNase1 structure using the Phyre program. Fig. 5 shows the
active and the allosteric sites in the predicted hASNase1 struc-
ture overlaid by the homologous regions of
Ec
ASNase1. Resi-
dues critical for catalysis in the
E. coli enzyme, such as Thr
14,
Thr
91, Asp
92,
and Lys
163,
overlay with high accuracy with those of
hASNase1, which has identical residues at these positions
(Thr
19, Thr
116, Asp
117,
and Lys
188) (Fig. 5A). Importantly, resi-

FIGURE 1. Alignment of the amino acid sequences of hASNase1 (UniProt accession number Q86U10), R. norvegicus (UniProt accession number O88820), E. coli (UniProt accession number P0A962), P. horikoshii (UniProt accession number O57797), and S. cerevisiae (UniProt accession number P38986) - asparaginases type I. The displayed sequence of hASNase1 corresponds to truncation 4, which showed the best expression pattern of the four
protein constructs produced in this work (see Fig. 2). Asterisks indicate the two threonine residues that are critical for L-asparaginase activity, and red arrows indicate residues that are located close to the allosteric sites. The alignment was performed using ClustalW (67), and the graph was generated using JalView (68). Blue shading indicates highly conserved amino acid residues.
dues Thr^{14} and Thr^{91} of the bacterial enzyme are considered to be the primary nucleophiles for the attack on the substrate L-asparagine (25). The presence of threonine residues at equivalent positions in the human enzyme is in favor of the view that the primary nucleophile in bacterial L-asparaginases (both cytoplasmic and periplasmic) is a threonine residue (47). This prompted us to investigate this assumption for hASNase1 by mutagenesis studies. Indeed, when point mutants T19A and T116A of hASNase1 were assayed, no L-asparaginase activity was detected (Table 1), thus indicating the critical catalytic role of these threonine residues in hASNase1.

Steady-state kinetic characterization of wild-type hASNase1 using L-Asn as substrate (there was no detectable L-glutaminase activity) revealed that the enzyme did not follow Michaelis-Menten kinetics. Instead, it exhibited a pronounced sigmoidal kinetic behavior, which is a hallmark of allosteric enzymes. The kinetic data were fitted using the Hill equation (Equation 1 under “Experimental Procedures”) from which we estimated a Hill coefficient (n_{H}) of 3.9 and an S_{0.5} value of 11.5 mM (Fig. 6). The modeled structure of the allosteric site is shown in Fig. 5B in direct comparison with that of EcASNase1. Significant differences between the two allosteric sites are seen in two key residues that directly interact with the substrate L-Asn: Glu^{266} and Ser^{327} in hASNase1 as opposed to Arg^{240} and Val^{302} in the bacterial enzyme. Aiming at further characterization of the allosteric sites of both enzymes, we generated a series of mutants and expressed, purified, and tested them for catalytic activity. Strikingly, hASNase1 tolerated none of the mutations introduced in the predicted allosteric site as evidenced by the total lack of activity of the mutant enzymes (Table 1). Given the high structural homology between hASNase1 and EcASNase1, we reasoned that the mutation E266R could potentially lower the S_{0.5} value of the human enzyme by mimicking the Arg^{240}}
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FIGURE 4. Size exclusion chromatography profiles upon separation using Superdex 200 column (GE Healthcare). A, standard gel filtration marker (Bio-Rad catalog number 151-1901). B, EcASNase1 chromatogram in 50 mM Tris-Cl, 100 mM NaCl, pH 8. C, profile of hASNase1 in the absence of L-Asn. Before sample loading, the column was equilibrated with 50 mM Tris-Cl, 100 mM NaCl, pH 8. D, hASNase1 in the presence of 20 mM L-Asn. The column was equilibrated with 50 mM Tris-Cl, 100 mM NaCl, 20 mM L-Asn, pH 8. In each case (B, C, and D), ~0.5 mg of enzyme was loaded on the column. mAU, milli-absorbance units.

FIGURE 5. A, modeled active site of hASNase1 overlaid by the active site of EcASNase1. Red highlights amino acids that are critical for activity of bacterial L-asparaginases: the catalytic triad formed by residues Thr116, Asp117, and Lys188 that ensure proper orientation of the substrate L-Asn (dark gray) and Thr14, which is responsible for subsequent nucleophilic attack (23, 69). Blue highlights the respective amino acids of hASNase1: Thr116, Asp117, Lys188, and Thr14. B, modeled allosteric site of hASNase1 (magenta residues) overlaid by the EcASNase1 allosteric site (blue residues). Key differences in homologous sites of the human enzyme are residues Glu206 versus Arg240 and Ser323 versus Val322. Other residues of the allosteric sites of the two enzymes are highly similar. Also shown is an L-Asn molecule (dark gray), which interacts with allosteric site residues. The interactions between L-Asn and the human enzyme were predicted based on the structure of the bacterial enzyme (Protein Data Bank code 2HIM).

TABLE 1

| Enzyme          | kcat | s0.5 | nH |
|-----------------|------|------|----|
| Wild-type hASNase1 | 6.7 ± 0.2 | 11.5 ± 0.8 | 3.9 ± 0.2 |
| hASNase1 T19A    | ND   | ND   | ND |
| hASNase1 T116A   | ND   | ND   | ND |
| hASNase1 E266R   | ND   | ND   | ND |
| hASNase1 E266S   | ND   | ND   | ND |
| hASNase1 T187A   | ND   | ND   | ND |
| hASNase1 T187S   | ND   | ND   | ND |
| hASNase1 C299S   | ND   | ND   | ND |
| Wild-type EcASNase1 | 7.4 ± 0.3 | 0.40 ± 0.05 | 3.5 ± 0.3 |
| EcASNase1 R240E  | 14.5 ± 0.5 | 1.8 ± 0.2 | 1.9 ± 0.2 |
| EcASNase1 C273S  | 16.8 ± 0.4 | 2.8 ± 0.3 | 2.7 ± 0.2 |

Kinetic data on wild-type and mutants of hASNase1 and EcASNase1

Steady-state kinetic analysis was performed by applying a continuous NADH-dependent assay (30). Enzyme concentrations were ~0.5 μM in a final volume of 1 ml of 50 mM Tris-Cl, 100 mM NaCl, pH 8 at 37 °C; the tested substrate concentrations covered the range of 0–10Km. Parameters were calculated from non-linear regression of a V/E versus [L-Asn] plot using the software SoftZymics (Igor Pro, WaveMetrics). ND, non-detectable; the limit of detection for the applied assays in the present study is 0.5 μM of ammonia produced in a 1-ml reaction volume. Data are shown as mean values ± S.D. of triplicate measurements.

site of the bacterial homolog. However, this mutation totally inactivated hASNase1. Other mutations introduced around the putative allosteric site of hASNase1, such as E266S, C299S, T187S, and T187A, also abolished activity (Table 1). Unlike the human enzyme, the E. coli enzyme tolerated mutations of its allosteric site. Mutation R240E resulted in an ~5-fold increase of S0.5 and 2-fold increase of kcat while simultaneously lowering the Hill coefficient nH from 3.55 to 1.9. The C273S mutation increased even further the S0.5 value (7-fold with respect to wild type), whereas the nH value slightly decreased (Fig. 6). All kinetic data of wild-type hASNase1 and EcASNase1 as well as of mutants generated in this work are summarized in Table 1. Taken together, these results suggest that despite the high similarity of sites predicted to be critical for their activities these two enzymes display different degrees of tolerance toward mutations of residues at these sites.

We additionally tested for potential hydrolysis of palmitoyl-lysoPC and lysoPI by hASNase1 and EcASNase1. We chose these two substrates because the rat 60-kDa lysophospholipase has been reported to hydrolyze palmitoyl-lysoPC (23), whereas lysoPI could serve as a substrate for the human homolog (24). However, we were unable to detect any activity of either hASNase1 or EcASNase1 using these two substrates under conditions where the control enzyme phospholipase A2 hydrolyzed L-α-phosphatidylcholine. The rate of the fluorescence decrease in the ADIFAB assay was the same for the blank (ADIFAB plus substrate without enzyme) and the sample (ADIFAB plus sub-
strate and enzyme) and accounted for ~10 arbitrary fluorescence units/min (data available upon request). This background activity is in line with another study (31), which also reported a constant slight fluorescence signal decrease in the absence of enzyme. In conclusion, our results demonstrate that the N-terminal domain of the 60-kDa full-length human lyso-
phospholipase can exist as a distinct folding unit that resembles the bacterial-type l-Asp-asparaginases, lacking the capacity to hydrolyze substrates other than l-Asn.

**Fluorescence Labeling of hASNase1**—Cysteine labeling allowed us to investigate the presence of an l-Asn binding site distinct from the catalytic site in the monomer. This additional binding site could play the role of an allosteric site of hASNase1, and its existence is supported not only by the sigmoidal kinetic behavior of enzyme described in the previous paragraph but also by our results obtained from substrate titration to a fluorescence-labeled, catalytically inactive version of the protein. Binding of the substrate l-Asn to the putative allosteric site was monitored by fluorescence signal changes of the Atto dye-labeled enzyme. The dye was covalently attached to cysteine residue Cys299, which according to the structural model built on the *E. coli* enzyme directly interacts with l-Asn bound to the allosteric site (Fig. 5B). The labeling of hASNase1 was confirmed by SDS-PAGE analysis as shown in Fig. 7 based on the observation that labeled proteins migrate more slowly than the unlabeled species (34). Fig. 8 shows a pronounced fluorescence signal decrease upon incubation of the catalytically inactive T19A mutant of hASNase1 with l-Asn in the concentration range of 0.1–40 mM. We did not label the wild-type enzyme because upon titration of l-Asn notably at high concentrations it would hydrolyze the substrate without allowing us to solely evaluate binding phenomena (the T19A mutant is assumed to maintain intact its allosteric site similarly to the wild-type enzyme). From the resulting binding curve, we estimated a $K_d$...
of 1 mM, which is about 10-fold lower than the $S_{0.5}$ (11 mM). Unlike T19A, the C299S mutant, which served as a negative control for labeling with the Atto dye, yielded a constant background fluorescence (due to labeling of other cysteines), showed no fluorescence signal changes upon l-Asn titration; constant background fluorescence results from labeling of other cysteines. The saturation binding curve for the T19A mutant was fitted to a hyperbolic function (Equation 2 in text) as described under “Experimental Procedures.” In the case of C299S, where no fluorescence change was observed, the points were fitted to a straight line. Error bars represent S.D. AFU, arbitrary fluorescence units.

**FIGURE 7.** SDS-PAGE analysis of unlabeled and labeled T19A and C299S mutants. Lane 1, unlabeled T19A; lane 2, labeled T19A; lane 3, unlabeled C299S; lane 4, labeled C299S; lane M, molecular mass markers. The electrophoretic pattern clearly shows the migration difference between the labeled and unlabeled enzyme species.

**FIGURE 8.** Plot of fluorescence signal change of Atto 465-labeled T19A and C299S HAsNase1 mutants as a function of l-Asn concentration. Shown are the fluorescence signal decreases in the case of T19A indicating an induced conformational change upon binding of l-Asn to either the allosteric or the active site. In contrast, the C299S mutant, which is not labeled at the putative allosteric site, showed no signal changes upon l-Asn titration; constant background fluorescence results from labeling of other cysteines. The saturation binding curve for the T19A mutant was fitted to a hyperbolic function (Equation 2 in text) as described under “Experimental Procedures.” In the case of C299S, where no fluorescence change was observed, the points were fitted to a straight line. Error bars represent S.D. AFU, arbitrary fluorescence units.

**FIGURE 9.** Temperature dependence of free energy change during unfolding of human and bacterial hASNase1. Shown is a plot of $\Delta G_{H2O}$ as obtained at various temperatures versus the respective temperatures for wild-type hASNase1 (△) and EcASNase1 (●). Data were fitted to the Gibbs-Helmholtz equation (Equation 3 in text) after determining $T_m$ and $\Delta H_m$ by thermal denaturation experiments (see “Experimental Procedures”). The standard deviation for $\Delta G_{H2O}$ (free energy change at zero concentration of urea), evaluated from three independent determinations on the wild-type enzymes at 25 °C was found to be 0.3 kcal/mol.

**TABLE 2** Thermodynamic parameters for wild-type hASNase1 and EcASNase1

| Enzyme | $\Delta C_p$ kcal/mol | $T_m$ °C | $\Delta H_m$ kcal/mol | $\Delta S_m$ cal/mol/K |
|--------|----------------------|---------|----------------------|----------------------|
| hASNase1 | 5.47 ± 0.20          | 58.40 ± 0.60 | 104.2 ± 1.5         | 314 ± 2              |
| EcASNase1 | 3.10 ± 0.15          | 59.50 ± 0.45 | 89.7 ± 0.9          | 270 ± 1.5           |

The $\Delta C_p$ values were calculated using the Gibbs-Helmholtz equation (see text) upon plotting the experimentally determined $\Delta G(T)$, $T_m$, $\Delta S_m$, and $\Delta H_m$ values. $\Delta S_m$, $T_m$, and $\Delta H_m$ were determined from linear regression analysis of the thermal denaturation of either enzyme. Parameters are represented as means ± S.D. of three measurements.

**Thermodynamic Characterization of hASNase1 and EcASNase1**—The free energy difference $\Delta G_{H2O}$ of folded and unfolded states, which is a measure of protein stability, was determined for both enzymes from urea denaturation experiments at different temperatures (37). Non-linear regression analysis was applied to the experimentally obtained $\Delta G_{H2O}$ values using the Gibbs-Helmholtz equation as a function of temperature, which allows the calculation of the $\Delta C_p$ parameter (48). Fig. 9 shows the stability curves for both enzymes studied in this work for direct comparison of their characteristic features under identical conditions. At 25 °C, the bacterial enzyme has a 2-fold higher $\Delta G_{H2O}$ in comparison with the human enzyme, although this difference appears to decline at higher temperatures (35 and 40 °C) as has also been observed in other proteins (49). Table 2 summarizes thermodynamic parameters obtained for hASNase1 and EcASNase1. Interestingly, the predicted $\Delta C_p$ values are considerably different for the two enzymes, possibly pointing to variations in the surface area that is exposed upon denaturation (50) and distinct mechanisms of enthalpy and entropy changes occurring upon unfolding (51). The two enzymes exhibited similar melting temperatures $T_m$ with hASNase1 showing higher $\Delta H_m$ and $\Delta S_m$ values, indicating higher enthalpy changes upon unfolding and a higher degree of disorder as compared with the E. coli enzyme (52).

**Differential Scanning Fluorimetry of hASNase1**—To analyze whether allosteric and active site mutations influenced the thermal stability of hASNase1, several of these hASNase1 mutants were studied by differential scanning fluorimetry. The melting curves were all monophasic (Fig. 10). The melting tem-
temperature ($T_m$) values are summarized in Table 3. Our results suggest that three point mutations (T19A, T187A, and T187S) had a significant stabilizing effect on the enzyme. In contrast, mutations T116A and C299S considerably lowered protein stability as compared with wild type, whereas the two mutations (E266S and E266R) introduced at the allosteric site Glu266 had almost no effect on the stability of hASNase1.

**TABLE 3**

Melting temperatures of wild-type and mutant hASNase1 as determined by differential scanning fluorometry

| Enzyme           | $T_m$ °C |
|------------------|----------|
| Wild-type hASNase1 | 51.3     |
| T19A             | 58.1     |
| T116A            | 41       |
| C299S            | 40       |
| E266S            | 52.5     |
| E266R            | 51       |
| T187A            | 57.9     |
| T187S            | 57.6     |

**DISCUSSION**

The present study focuses on the biochemical analysis of the N-terminal domain of a human 60-kDa protein designated as lysophospholipase. The full-length version of this two-do-
main protein has been only poorly characterized in one report (24). Our primary motivation to produce and functionally characterize the N-terminal domain of this protein, which structurally and according to its catalytic in vitro properties significantly resembles E. coli cytoplasmic l-asparaginase (ansA; EcASNase1), originated from our previous work on human enzymes that possess l-asparaginase activity (10, 19, 21). The discovery, molecular engineering, and in vitro evolution of catalytically efficient human l-asparaginas are thought to lay the basis for the replacement of bacterial l-asparaginas presently used as antileukemia therapeutics despite adverse side effects mainly attributed to their bacterial origins (54).

An early study on the rat 60-kDa lysophospholipase (23) assigned three distinct activities to this enzyme acting as lysophospholipase, l-asparaginase, and acetylhydrolase on the platelet-activating factor. Several years later, a report on the human 60-kDa lysophospholipase (24) revealed a specific role of this enzyme as an interacting partner of the serum- and glucocorticoid-induced serine/threonine protein kinase Sgk1, an enzyme involved in various cell proliferation pathways. In contrast to its rat homolog, the human enzyme was not reported to bear l-asparaginase activity (24). However, in both these studies, the distinct enzymatic activities were determined only qualitatively using lysates from cells overexpressing the respective genes rather than purified enzyme preparations.

Here, we show that the N-terminal l-asparaginase domain (residues 1–369) of the 573-residue human protein, that we termed hASNase1, can be produced in soluble form in E. coli. This protein showed l-asparaginase activity with a maximum catalytic rate of about 7 s⁻¹, which is very similar to that of EcASNase1, implying that this N-terminal domain forms a stable and functional folding unit in the absence of the C-terminal putative ankyrin repeat domain. Kinetic and mutational characterization of hASNase1 revealed strong positive allosteric modulation in the velocity versus substrate plot similar to its E. coli homolog. To validate our experimental strategy for characterizing this human counterpart of the cytosolic E. coli l-asparaginase, we also recombinantly produced the bacterial protein to directly compare the two enzymes under identical assay conditions. Our kinetic data obtained on the bacterial enzyme are consistent with a previous study that centered on its structural analysis and allosteric regulation (25).

Given the pronounced sigmoidal kinetic behavior that we observed in steady-state kinetics of l-asparagine hydrolysis, we hypothesized that hASNase1 possesses an allosteric site that could act as “sensor” for the presence of substrate. Our dye labeling approach targeting the Cys²⁵⁹ residue indicated fluorescence signal changes upon binding of l-Asn, thus strengthening the idea of the existence of such an allosteric site that can be regulated by the substrate l-Asn. In the absence of information on the three-dimensional structure of hASNase1, which could provide more detailed insight into the catalytic mechanism and allosteric regulation of the enzyme, we referred to a structural homology-model based on the EcASNase1 crystal structure, which allowed us to define the active site and an allosteric site of the human enzyme. The S₀.₅ value of 11.5 mM determined for hASNase1 is about 10-fold higher than that reported for the E. coli enzyme, raising questions about the physiological role of the full-length protein and its N-terminal domain displaying l-asparaginase activity as shown in this work. The relatively high S₀.₅ value determined for the human enzyme falls well within the range of free concentrations of intracellular amino acids (>10 mM), including l-Asn, reported for mammalian cells (55, 56), thus making the enzyme operate efficiently, particularly at elevated substrate concentrations.

The homo-oligomeric EcASNase1 displays sigmoidal kinetics, which can be explained by positive cooperativity induced by the substrate l-Asn. In contrast, hASNase1, which shows non-Michaelis-Menten kinetics similar to its E. coli homolog, is monomeric under the conditions of activity measurements. In fact, a number of monomeric enzymes exhibiting allosteric behavior have been reported of which the most thoroughly characterized is human glucokinase (57, 58). However, glucokinase (also called hexokinase IV), unlike l-asparaginase, is a two-substrate (ATP plus glucose) enzyme and displays a moderate degree of alloster (59). However, monomeric enzymes with single binding sites, like hASNase1, can also show alloster (60).

Two basic models have been put forward aiming at the mechanistic elaboration of monomeric alloster: the mnemonic model (61) and the ligand-induced slow transition model (62). Both models assume the existence of two different enzyme conformations that are characterized by distinct affinities for the substrate (low and high affinity states). Depending on the substrate concentration, certain conformational changes may occur, thereby perturbing the equilibrium of the two states in favor of the high affinity state and an increased catalytic activity. Based on the observed sigmoidal kinetic behavior and the monomeric state of hASNase1, it appears plausible to assume that this is another example of a monomeric enzyme exhibiting positive allosteric regulation. Importantly, given the fact that l-Asn plays a dual role of being both substrate and regulator of hASNase1, l-Asn can be considered as a homotropic allosteric effector of this enzyme, adding to the steadily growing number of allosterically regulated proteins (63).

The role of the putative ankyrin repeat structure located in the C-terminal part of the full-length protein remains unknown at present. In numerous other proteins, ankyrin repeats were shown to mediate protein-protein interactions, and therefore, they may play crucial roles in cellular signaling events (64–66). The study of the kinetics of the full-length 60-kDa human lysophospholipase could provide information on the potential influence of the ankyrin repeat domain on the l-asparaginase activity predicted to reside in the N-terminal part. On the other hand, when we tested palmitoyl-lysoPC and lysoPI as potential substrates for hASNase1, we detected no lipase activity. Our biochemical data clearly support the view that the N-terminal domain bears l-asparaginase activity that exhibits positive allosteric regulation by the substrate l-Asn. A particular characteristic of hASNase1 is its monomeric state in conjunction with its pronounced sigmoidal steady-state kinetics. We would like to emphasize that the distinct enzymatic activities assigned to rat and human 60-kDa lysophospholipases warrant further analyses at the cellular level to elucidate the physiological role of these two-domain proteins. Our studies on l-asparaginase activity inherent to the N-terminal domain of this 60-kDa protein expands the basis of our work aiming at the identification
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and molecular engineering of enzymes of human origin (10, 21) that might become suitable for replacing bacterial enzymes as approved therapeutics in the treatment of leukemias.

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