Purification, characterization, cytotoxicity and anticancer activities of L-asparaginase, anti-colon cancer protein, from the newly isolated alkaliphilic Streptomyces fradiae NEAE-82

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L-asparaginase is an important enzyme as therapeutic agents used in combination with other drugs in the treatment of acute lymphoblastic leukemia. A newly isolated actinomycetes strain, Streptomyces sp. NEAE-82, was potentially producing extracellular L-asparaginase, it was identified as Streptomyces fradiae NEAE-82, sequencing product was deposited in the GenBank database under accession number KJ467538. L-asparaginase was purified from the crude enzyme using ammonium sulfate precipitation, dialysis and ion exchange chromatography using DEAE Sepharose CL-6B. Further the kinetic studies of purified enzyme were carried out. The optimum pH, temperature and incubation time for maximum L-asparaginase activity were found to be 8.5, 40 °C and 30 min, respectively. The optimum substrate concentration was found to be 0.06 M. The Km and Vmax of the enzyme were 0.01007 M and 95.08 U/ml min−1, respectively. The half-life time (T½) was 184.91 min at 50 °C, while being 179.53 min at 60 °C. The molecular weight of the subunits of L-asparaginase was found to be approximately 53 kDa by SDS–PAGE analysis. The purified L-asparaginase showed a final specific activity of 30.636 U/mg protein and was purified 3.338-fold. The present work for the first time reported more information in the production, purification and characterization of L-asparaginase produced by newly isolated actinomycetes Streptomyces fradiae NEAE-82.

L-asparaginase (L-asparagine aminohydrolase, E.C. 3.5.1.1) is the drug of choice used in combination therapy with other drugs in the treatment of acute lymphoblastic leukemia chemotherapy in children1. The demand for L-asparaginase will increase several fold in coming years due to its potential industrial applications as food processing aid in addition to its clinical applications2. L-asparaginase is highlighted as a key drug in the treatment of extranodal NK/T-cell lymphoma3.

Purification of a protein is an important step for characterization of its physical and biological properties. Moreover, for effective therapeutic use of a protein, it must be free of any contaminants and impurities. However, clinical employments of L-asparaginase are accompanied with fatal allergenic reactions to the patients4. These effects are mainly due to L-asparaginase associated L-glutaminase activity and bacterial endotoxins in enzyme preparations5. Several research groups have studied L-asparaginase production and purification in attempt to

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minimize impurities that produce allergenic reactions. The L-asparaginase enzyme was purified from *Penicillium* sp. that was grown on submerged fermentation. Different purification steps including salt precipitation, followed by separation on sephadex G-100-120 gel filtration and DEAE to obtain pure enzyme preparation. The purified enzyme showed 13.97 IU/mg specific activity. The polyacrylamide gel electrophoresis of the pure enzyme exhibited one protein of 66 kDa. The enzyme showed maximum activity at 7.0 pH and 37 °C and *Km* value 4 × 10⁻³ M⁻¹.

Three marine soil isolates (S3, S4 and K8) synthesized asparaginase with yield ranging between 24.6 to 49.2 IU/ml. Isolate S3 showed the highest productivity of 49.2 IU/ml and optimum activity at pH 7.5 and 50°C. The *Km* and Vmax of partially purified enzyme were approximately 24µM and 51 IU/ml, respectively.

Furthermore, the physiological and biochemical reactions of strain NEAE-82 are shown in Table 2. It exhibited no antimicrobial activities against *Klebsiella pneumoniae*, *Rhizoctonia solani*, *Fusarium oxysporum*, *Aspergillus niger* in Table 2. It exhibited no antimicrobial activities against *Klebsiella pneumoniae*, *Rhizoctonia solani*, *Fusarium oxysporum*, *Aspergillus niger*

### Results and Discussion

#### Morphology and cultural characteristics of the isolate no. NEAE-82.

The colonial morphology of a 14 day culture of strain NEAE-82 grown on yeast extract/malt extract agar (ISP 2 medium) revealed that strain NEAE-82 had the typical characteristics of the genus *Streptomyces*²¹. It is aerobic, mesophilic, Gram-positive actinomycete that develops abundant and well-developed substrate and aerial mycelium. Strain NEAE-82 produced reddish brown aerial mycelium (Fig. 1) on yeast extract-malt extract agar, oatmeal agar, inorganic salt-starch agar, and peptone-yeast extract iron agar. Not-distinctive aerial mycelium on glycerol-asparagine agar and tyrosine agar. Substrate mycelium with no distinctive pigments on most tested medium and brown substrate mycelium was produced on peptone-yeast extract iron agar (Table 1). Substrate mycelium pigment is not a pH indicator. No pigment found in medium in yeast extract -malt extract agar, oatmeal agar inorganic salt-starch agar, glycerol–asparagine agar or tyrosine agar, faint brown pigments formed in peptone-yeast extract iron agar. Melanoid pigments not formed in peptone-yeast extract iron agar and tyrosine agar. Strain NEAE-82 grew well on yeast extract -malt extract agar (ISP medium 2), oatmeal agar (ISP medium 3), inorganic salt-starch agar (ISP medium 4), peptone-yeast extract iron agar (ISP medium 6) but poor growth on glycerol–asparagine agar (ISP medium 5) and tyrosine agar (ISP medium 7). Strain NEAE-82 formed an extensively branched substrate mycelium and aerial hyphae which differentiated into long straight spore chains. Spore chains in section Retinaculapierti including open spiral spore chains (Fig. 2). Flexuous or spiral spore chains are seen on starch-nitrate agar. Mature spore chains generally have 10–50 spores per chain; longer chains are sometimes observed. Spore surface is smooth. Verticils are not present. The mycelium does not fragment.

#### Physiological characteristics.

The physiological and biochemical reactions of strain NEAE-82 are shown in Table 2. It exhibited no antimicrobial activities against *Staphylococcus aureus*, *Alternaria solani* and *Bipolaris oryzae*. *Saccharomyces cerevisiae*, *Candida albicans*, *Bacillus subtilis*, *Escherichia coli*, *Pseudomonas aeruginosa*, *Rhizoctonia solani*, *Fusarium oxysporum*, *Aspergillus niger* and *Klebsiella pneumoniae*. α-amylose (starch hydrolysis) (Fig. 3), protease (degradation of casein), cellulase (growth on cellulose), chitosanase and L-asparaginase of strain NEAE-82 were produced. Melanin production, lecithinase activity and uricase were not produced. Coagulation and peptonization of milk (Fig. 3) and gelatin liquefaction were positive. Maximum NaCl tolerance falls into a clade together with *Streptomyces somaliensis* strain DSM 40738 (GenBank/EMBL/DDBJ accession No. NR_025292.1) and *Streptomyces fradiae* strain CBT BR13 (GenBank/EMBL/DDBJ) accession No. KP230701.1).

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**16S rRNA gene sequence comparisons and phylogenetic analysis.** The 16S rRNA gene sequence (1508 bp) was determined for strain NEAE-82. A BLAST search⁴ of the GenBank database using this sequence showed its similarity to that of many species of the genus *Streptomyces*. A phylogenetic tree (Fig. 4) based on 16S rRNA gene sequences of members of the genus *Streptomyces* was constructed according to the neighbour-joining method of Saitou and Nei⁷ with MEGA⁶²⁰. This tree shows the close phylogenetic association of strain NEAE-82 with certain other *Streptomyces* species. Phylogenetic analysis indicated that the strain NEAE-82 consistently falls into a clade together with *Streptomyces somaliensis* strain DSM 40738 (GenBank/EMBL/DDBJ accession No. NR_025292.1) and *Streptomyces fradiae* strain CBT BR13 (GenBank/EMBL/DDBJ) accession No. KP230701.1).
Figure 1. Color of the aerial mycelium of *Streptomyces* sp. NEAE-82 grown on starch-nitrate agar medium for 7–14 days of incubation at 30 °C.

| Medium                        | Aerial mycelium (spore-colour en masse) | Substrate mycelium | Diffusible pigment | Growth   |
|-------------------------------|----------------------------------------|--------------------|--------------------|----------|
| ISP medium 2 (Yeast extract-malt extract agar) | Reddish brown                         | Not-distinctive   | Non-pigmented      | Excellent |
| ISP medium 3 (Oatmeal agar)   | Reddish brown                         | Not-distinctive   | Non-pigmented      | Excellent |
| ISP medium 4 (Inorganic salt-starch agar) | Reddish brown                         | Not-distinctive   | Non-pigmented      | Excellent |
| ISP medium 5 (Glycerol asparagines agar) | Not-distinctive                      | Not-distinctive   | Non-pigmented      | Weak     |
| ISP medium 6 (Peptone-yeast extract iron agar) | Reddish brown                        | Brown             | Faint brown        | Excellent |
| ISP medium 7 (Tyrosine agar)  | Not-distinctive                        | Not-distinctive   | Non-pigmented      | Weak     |

Table 1. Culture characteristics of the *Streptomyces* sp. strain NEAE-82. The substrate mycelium pigment was not pH sensitive when tested with 0.05 N NaOH or 0.05 N HCl. The diffusible pigment was not pH sensitive when tested with 0.05 N NaOH or 0.05 N HCl.

Figure 2. Scanning electron micrograph showing the spore-chain morphology and spore-surface ornamentation of strain NEAE-82 grown on starch nitrate agar medium for 14 days at 30 °C at magnification of 4000X (A) and 10000X (B).
On the basis of the collected data and in view of the comparative study of the recorded properties of isolate No. NEAE-82 (Table 2) in relation to the closest related species of the genus *Streptomyces*, it is most closely related to the type strains of *Streptomyces fradiae* strain CBT BR13 (GenBank/EMBL/DDBJ accession No. KP230701.1) (99% sequence similarity)\(^2^1\). Therefore, this strain was identified as *Streptomyces fradiae* strain NEAE-82 and its sequencing product was deposited in the GenBank database under accession number KJ467538.

### Purification of L-asparaginase from *Streptomyces fradiae* NEAE-82

The crude culture filtrate of *Streptomyces fradiae* NEAE-82 had a total activity of 29507.688 U with protein content 3214.705 mg; the specific activity was 9.179 U/mg protein. The ammonium sulphate concentrated enzyme preparation had a protein content of 196.6 mg and the specific activity was 13.008 U/mg protein, showing purification fold of 1.417 with enzyme recovery at this step was 8.667 per cent. The fractions collected after ammonium sulphate precipitation were loaded on the column packed with DEAE Sepharose CL-6B and fractions of 2 ml were collected and analyzed for enzyme activity and protein content (Fig. 5). A total of 285 fractions were collected that showed one major L-asparaginase peak on the chromatogram. The total activity was 1442.393 with 47.081 mg protein after ion exchange chromatography through DEAE Sephacel CL-6B column. The specific activity of the purified enzyme was 30.636 U/mg of protein. Summary of the purification steps of the L-asparaginase produced by *Streptomyces fradiae* NEAE-82 is presented in Table 3.

### Kinetics properties of the purified L-asparaginase

The activity of L-asparaginase of *Streptomyces fradiae* NEAE-82 was evaluated at different levels of pH, temperature, effect of substrate concentration and incubation time.

![Table 2: Phenotypic properties that separate strain *Streptomyces NEAE-82* from related *Streptomyces* species.](https://www.nature.com/scientificreports/)
Effect of pH on L-asparaginase activity. L-asparaginase activity was studied as a function of pH in the range between 4.5–10.5 (Fig. 6). L-asparaginase was active over a broad pH range (4.5–10.5). The enzyme activity increased gradually till pH 8.5 with maximum activity 20.932 IU (relative activity, 100%). At higher pHs, enzyme activity decreased.
activity was decreased. The enzyme retains up to 29.348% of activity at pH 10.5 compared to 44.021% at pH 4.5. It is known that L-asparaginase can completely lose its activity and also recover it partially, depending on the exposure conditions. These results coincide with that of Dhevagi and Poorani\textsuperscript{23} who reported the maximal activity of L-asparaginase as a function of the pH of the reaction.

| Purification step                  | Total protein content (mg) | Total activity (U) | Specific activity (U/mg protein) | Recovery (%) | Purification fold |
|-----------------------------------|-----------------------------|--------------------|---------------------------------|--------------|-------------------|
| Culture filtrate                  | 3214.705                    | 29507.688          | 9.179                           | 100          | 1                 |
| (NH4)\textsubscript{2}SO\textsubscript{4} post dialysis | 196.6                       | 2557.325           | 13.008                          | 8.667        | 1.417             |
| Ion exchange on DEAE Sepharose CL-6B | 47.081                     | 1442.393           | 30.636                          | 4.888        | 3.338             |

Table 3. Summary of the purification steps of the L-asparaginase produced by \textit{Streptomyces fradiae} NEAE-82.
L-asparaginase activity of *Streptomyces* sp. PDK7 was between pH 8.0 and 8.5, and the optimal L-asparaginase activity extracted from *Streptomyces gulbargensis* was 9.0. L-asparaginase is one of the amidases that are generally active and stable at neutral and alkaline pH, whereas, pH 5.0 to 9.0 were reported earlier to be optimum for amidase activity. L-asparaginase, purified from *Streptomyces acrimycini* NGP, exhibited maximum activity at pH 7.0; membrane bound L-asparaginase from *Tetrahymena pyriformis* acts optimally at pH 9.6 and the optimal L-asparaginase activity from *Corynebacterium glutamicum* was reported at pH 7.0. Maximum activity of purified L-asparaginase occurred at pH 9 was obtained for *Pseudomonas aeruginosa* 5007122.

**Effect of temperature on L-asparaginase activity.** The temperature optimum of L-asparaginase from *Streptomyces fradiae* NEAE-82 is shown in Fig. 7. It was active at wide range of temperature condition from 25–60 °C. The maximum L-asparaginase activity of 26.848 IU was obtained at 40 °C. At higher temperature the L-asparaginase activity declined. The enzyme retains 50.85% of its activity at 60 °C. Our results were in agreement with a previous study which reported that the maximum activity of L-asparaginase purified from *Streptomyces gulbargensis* was at 40 °C11. Manna et al.28 have found 37 °C to be the optimum temperature for the enzyme activity obtained from *Pseudomonas stutzeri* MB-405. However, optimum temperature for L-asparaginase activity obtained from *Erwinia* sp. showed maximum activity at 35 °C29.

Figure 7. Effect of the temperature on L-asparaginase activity.

Figure 8. Effect of the substrate concentration of the reaction on L-asparaginase activity.
Effect of substrate concentration on the activity of L-asparaginase. In this experiment, the influence of substrate concentration on L-asparaginase activity was examined by using different concentration of substrate ranging from 0.02 to 0.1 Molar to determine the optimum concentration of substrate required to give the highest L-asparaginase activity. The results in Fig. 8 showed a gradual increase in the enzyme activity with the increase in substrate concentration from 0.02 to 0.06 Molar. However, further increase in substrate concentration (0.07–0.1 Molar) lead to decrease in enzyme activity to 53.03% with 0.1 Molar substrate. The optimum substrate concentration for L-asparaginase activity was observed at 0.06 Molar.

The typical Michaelis-Menten relationship was obtained between the substrate concentrations and the initial velocity of the reaction. Michaelis-Menten plot showed in Fig. 9 illustrated the $K_m$ and $V_{max}$ values for L-asparaginase enzyme. The plot gave $K_m$ value of 0.01007 M and $V_{max}$ of 95.08 U ml$^{-1}$ min$^{-1}$ for the hydrolysis of L-asparagine. $K_m$ value is defined as the substrate concentrations that result in half maximal velocity for the enzymatic reaction, or an equivalent way of stating substrate concentration at which half of the enzyme active sites in the sample are filled (i.e. saturated) by substrate molecules in the steady state. It can be used as a relative measure of substrate affinity with the studied enzyme. $K_m$ reflects the affinity of the enzyme for its substrate$^{30}$. The lower the $K_m$ is the stronger binding ability the enzyme has. $V_{max}$ is the limiting velocity as substrate concentrations get very large. $V_{max}$ is expressed in units of product formed per unit of time. If the molar concentration of enzyme is known, $V_{max}$ be expressed as moles of product formed per second per mole of enzyme sites$^{31}$. This is the turnover
number, the number of molecules of substrate converted to product by one enzyme site per second. However, there are many factors that affect on kinetic parameters of the enzymes ($K_m$ and $V_{max}$) such as; type of enzyme, different forms of enzyme (crude, modified or purified), changes in enzyme conditions (pH, temperature, etc.), source of the enzyme (different microorganisms), type of used substrates, and the assay procedures32.

Effect of incubation time on enzyme activity. The L-asparaginase activity (Fig. 10), increased as the incubation time increased up to 30 min (L-asparaginase activity of 34.128 IU). After which only a slight decrease in L-asparaginase activity was observed. El-Bessoumy et al.22 reported that, maximum activity of L-asparaginase purified from *Pseudomonas aeruginosa* 50071 was at 30 min. In addition, the effect of incubation time on the activity of purified L-asparaginase from *Streptomyces noursei* showed that the activity reached its maximum at 35 min15. The decrease in L-asparaginase activity was observed after longer period of incubation with substrate. The decrease may due to the product inhibition.

| Thermal inactivation rate constant ($k_d$ (min$^{-1}$))$^*$ | 50 °C | 0.0037 |
|-------------------------------------------------------------|-------|--------|
| 60 °C                                                       | 0.0039 |
| 70 °C                                                       | 0.0046 |
| 80 °C                                                       | 0.0092 |

| Thermal inactivation rate constant ($k_d$ 2(min$^{-1}$))$^{**}$ | 50 °C | 0.0021 |
|-----------------------------------------------------------------|-------|--------|
| 60 °C                                                           | 0.0021 |
| 70 °C                                                           | 0.0026 |
| 80 °C                                                           | 0.0051 |

Table 4.  Half life time ($T_{1/2}$) and heat deactivation constant ($k_d$) of L-asparaginase produced by *Streptomyces fradiae* NEAE-82. $^*$ $k_d$ is the deactivation constant after losing 50% of initial activity (at $T_{1/2}$). $^{**}$ $k_d$ 2 is the deactivation constant after losing 90% of initial activity (at D value).
Thermal stability. The effect of temperature on the stability of L-asparaginase showed maximum enzyme activity at 50 °C (Fig. 11). Around 86.126% of the initial activity was retained by the enzyme after 20 min of incubation at 50 °C. About 72.072% of L-asparaginase activity was lost after incubation at 50 °C for 90 min, while a rapid decrease in the enzyme activity (16.874%) was observed after incubation at 80 °C for 90 min. Thermal inactivation process of the enzyme follows the theoretical curve of a sample first order reaction. However, linear regression of the obtained data was assayed to determine half life time (T1/2) as shown in Table 4. The half-life time (T1/2) was 184.91 min at 50 °С, while being 179.53 min at 60 °С. On the other hand, destruction of enzyme activity was observed at 80 °C with low half-life time (75.36 min). It can be concluded from the previous results that the higher thermal stability behavior of L-asparaginase was at 50 °C. Results presented in Table 4 were normalized to activity of un-incubated enzyme (time 0 = 100%) for the percentage of activity remaining. Heat inactivation half-life (T1/2) and heat deactivation constant (k) were determined by fitting the data to a first-order decay curve using Graph-Pad Prism software. An earlier study reported no significant loss of L-asparaginase activity purified from Streptomyces radiopugnans MS1, when the enzyme was pre-incubated at 40 °C for 60 min33. Similar results were recorded with Streptomyces noursei15, Erwinia carotovora34, Pseudomonas stutzeri MB 40528.

A source-dependent variation of physicochemical and biochemical properties, like optimum pH, temperature, substrate specificity, inhibition pattern, etc., of microbial enzymes is well documented8. Comparative evaluation of L-asparaginase for its potential activity from different microbial sources revealed that biochemical and therapeutic properties differ with source of strain in addition to enzyme properties. L-asparaginases from Erwinia chrysanthemi and E. coli are currently in clinical use as effective drugs in the treatment of the acute lymphoblastic leukemia. Comparison of the two enzymes, lead to the conclusion that most patients allergic to the E. coli- L-asparaginase, Erwinia asparaginase is considered less toxic and is frequently employed compared with allergic reactions to E. coli- L-asparaginase. However, Erwinia asparaginase had a shorter half life than E. coli4. The optimum temperature for L-asparaginase activity obtained from E. coli showed maximum activity at 37 °C and pH 7–835. The maximum L-asparaginase activity obtained from Erwinia aroidae46 occurred between pH 7 and 8 and optimum temperature for L-asparaginase activity obtained from Erwinia sp. showed maximum activity...
at 35 °C. However, the characterization of the enzyme produced by newly isolated actinomycetes *Streptomyces fradiae* NEAE-82 revealed that the optimum pH, temperature and incubation time for maximum L-asparaginase activity were found to be 8.5, 40 °C and 30 min, respectively. It was active at wide range of temperature condition from 25–60 °C and active over broad pH ranges (4.5–10.5). The physiological pH is one of the perquisites for anti-tumor activity under alkaline pH condition, L-asparaginase becomes a competitive inhibitor. This property of the enzyme is most suitable for complete elimination of asparagines from the body when tumor patient is treated.

Figure 14. The anticancer effects of the purified L-asparaginase on Caco2, Hep2 and HepG2 cells.

Figure 15. The anticancer effect of the purified L-asparaginase on Caco2 cells after 48 of treatment, cells undergoing apoptosis are characterized by cellular rounding up, shrinkage, membrane blebbing and loss of cell adhesion.
with L-asparaginase in-vivo and clarified that the enzyme produced by *Streptomyces fradiae* NEAE-82 under the present study (optimum pH 8.5) has effective antitumor activity.

**Molecular weight determination by Sodium Dodecyl Sulphate- Polyacrylamide Gel Electrophoresis (SDS–PAGE).** The molecular weight of the extracted enzyme was determined by performing SDS-PAGE (sodium dodecyl sulfate–polyacrylamide gel electrophoresis) according to the method of Laemmli28, with a separating acrylamide gel of 10% and stacking gel 5% containing 0.1% SDS. After the electrophoresis, the gel was stained in 0.025 Coomassie brilliant blue R-250 and destained with a solution of methanol-acetic acid and water in the ratio of 4:1:5. The molecular weight of the purified L-asparaginase was determined in comparison with standard molecular weight markers (molecular mass range: 9–178 kDa). SDS–PAGE of the enzyme preparation revealed only a single distinctive protein band for the pure preparation of L-asparaginase with an apparent molecular weight of 53 kDa (Fig. 12).

L-asparaginase is known as a homotetramer. Four active sites are located at the interface between two subunits forming an intimate dimer, and hence the asparaginase is more accurately described as a dimer of dimers with a molecular mass of approximately 120 to 160 kDa possessing antitumor activity46. The molecular weight of the L-asparaginase was found to be varied according to the source of enzyme like 80 kDa in *E. coli*11, 140 kDa in *Streptomyces* sp. PDK23 and 116 kDa in *S. albidoflavus*. Purified L-asparaginase from *S. tendae* checked by SDS-PAGE revealed a distinct protein band near 97.4 kDa41. In this respect, L-asparaginases purified from *Pseudomonas stutzeri* MB-405, *Thermus thermophiles* and *Escherichia coli* were with smaller molecular weight values ranging from 33–34 kDa28,42. Other reports on production and purification of L-asparaginase from *E. coli* revealed that the molecular weight was determined as 153 kDa with the help of SDS-PAGE43. The functional L-asparaginase from *E. coli* is a homotetramer with a molecular weight of about 142 kDa44. Purified L-asparaginase from *Streptomyces gulbargensis*31, *Streptomyces albidoflavus*, *Streptomyces PDK23* and *Streptomyces noursei*45 exhibited a molecular weight of 85, 112, 140 kDa and 102 kDa, respectively. Reports on production and purification of L-asparaginase from *Pseudomonas aeruginosa* 50071 by SDS–PAGE revealed a peptide chain with molecular weight of 160 kDa43.

**Cytotoxicity and anticancer activities of L-asparaginase on non-cancerous and cancer cells.** The safety pattern of purified L-asparaginase was scanned on human fibroblast cells. Generally, the obtained data revealed that, the treatment IC50 on all cells ranged from 2 to 4 U/ml (Fig. 13). The anti-proliferative activity of L-asparaginase on cancer cells was quantitatively estimated on HepG2, Hep2 and Caco2 cells. The obtained data generally indicated that, the activity of the extract against Caco2 was superior to that with either HepG2 or Hep2 cells (Figs 14 and 15, Table 5). Up on cancer cells treatment, the selected recommended dose (4 U/ml) showed anti-proliferation activities against Caco2 cells with percentages 80.9 with cancer cell selectivity index reached 4. On the other hand, the anticancer activity of the purified on Hep2 and Caco2 cells reached 39.95 and 36.79%, respectively with selectivity index 0.88 and 0.64, respectively. Sialyl Lewis X (sLex) is a tetrasaccharide carbohydrate that is often linked to cell-surface glycoproteins. Sialic acids in general play important roles in biological characteristics of cancer and other cells because it overexpressed in some tumor cell types and that are implicated in cellular invasiveness, differentiation and tumourigenecity45,46. Sialyl Lea (sLea) is a tetrasaccharide carbohydrate that is often linked to cell-surface glycoproteins. Sialic acids in general play important roles in biological characteristics of cancer and other cells because it overexpressed in some tumor cell types and that are implicated in cellular invasiveness, differentiation and tumourigenecity45,46.

**Materials and Methods**

**Microorganisms and cultural conditions.** *Streptomyces* spp. used in this study were isolated from various soil samples collected from different localities of Egypt and Saudi Arabia. Actinomycetes had been isolated from the soil using standard dilution plate method procedure on Petri plates containing starch nitrate agar medium of the following composition (g/L): Starch, 20; KNO3, 2; K2HPO4, 1; MgSO4.7H2O, 0.5; NaCl, 0.5; CaCO3, 3; FeSO4.7H2O, 0.01; agar, 20 and distilled water up to 1 L; then plates were incubated for a period of 7 days at 30 °C. *Streptomyces* isolates were purified and maintained as spore suspensions in 20% (v/v) glycerol at −20°C for subsequent investigation.

**Screening of L-asparaginase production by plate assay.** It is generally observed that L-asparaginase production is accompanied by an increase in pH of the culture filtrates29. The plate assay was based on Gulati et al.31 method with the incorporation of pH indicator phenol red (prepared in ethanol) in medium containing L-asparaginase (sole nitrogen source). Phenol red at acidic pH is yellow and at alkaline pH turns pink, thus a pink zone is formed around microbial colonies producing L-asparaginase. Screening of potential L-asparaginase

| Cell type | Cancer cells selectivity index (SI) |
|-----------|------------------------------------|
| HepG2 cells | 0.6472                             |
| Caco2 cells | 4                                 |
| Hep2 cells  | 0.8843                             |

Table 5. The Cancer cell selectivity index of the purified L-asparaginase.
producing actinomycetes was carried out with the use of asparagine dextrose salts agar (ADS Agar) (asparagine 1.0%, dextrose 0.2%, K$_2$HPO$_4$ 0.1%, MgSO$_4$ 0.05%, agar 1.5%), pH was adjusted to 6.8 and supplemented with phenol red as a pH indicator (0.009% final concentration) and sterilized at 1.5 atmospheric pressure for 20 min. Inoculated plates were incubated at 30°C for 7 days. Plates were examined for change in color of medium from yellowish to pink due to change of pH indicating the positive asparaginase activity. Colonies with pink zones were considered as L-asparaginase-producing strains. Isolates exhibiting L-asparaginase activity were selected for further study. Control plates were prepared as uninoculated medium and medium without dye.

**Inoculum preparation.** 250 ml Erlenmeyer flasks containing 50 ml of asparagine dextrose salts broth (L-asparagine 1.0%, dextrose 0.2%, K$_2$HPO$_4$ 0.1%, MgSO$_4$ 0.05%) were inoculated with three disks of 8 mm diameter taken from the 7 days old stock culture grown on starch nagar agar medium. The flasks were incubated for 48–72 h in a rotatory incubator shaker at 30°C and 150 rpm and were used as inoculum for subsequent experiments.

**Production of L-asparaginase by submerged fermentation.** The selected strain was cultured in fifty ml of asparagine dextrose salts broth medium (at a specified pH) dispersed in 250 ml Erlenmeyer conical flasks. The inoculated flasks were incubated on a rotatory incubator shaker at 30–37°C with shaking at 150–250 rpm. After the specified incubation time for each set of experimental trials, the mycelium of the tested isolate was collected by centrifugation at 5000 × g for 30 min at 4°C.

**Assay of L-asparaginase activity.** L-asparaginase activity was determined by measuring the amount of ammonia released by nesslerization according to the method described by Wriston and Yellin. The reaction mixture containing 1.5 ml of 0.04 M L-asparaginase prepared in 0.05 M Tris-HCl buffer, pH 8.6 and 0.5 ml of an enzyme to make up the total volume to 2 ml. The tubes were incubated at 37°C for 30 minutes. The reaction was stopped by adding 0.5 ml of 1.5 M trichloroacetic acid (TCA). The blank was prepared by adding enzyme after the addition of TCA. The precipitated protein was removed by centrifugation at 10,000 × g for 5 min and the liberated ammonia in the supernatant was determined colorimetrically by direct nesslerization by adding 1 ml Nessler’s reagent into tubes containing 0.5 ml of clear supernatant and 7 ml distilled water and incubated at room temperature for 20 min. A yellow coloration indicates the presence of ammonia: at higher concentrations, a brown precipitate may form. The yellow color was read using a UV-visible spectrophotometer (Optizen Pop –UV/Vis spectrophotometer) at 480 nm. The amount of ammonia liberated was calculated using ammonium (ammonium chloride) standard curve. One unit (U) of L-asparaginase is defined as the amount of enzyme which catalyzed the formation of 1 μmole of ammonia from L-asparagine per minute at 37°C and pH 8.6. The enzyme activity was expressed in terms of units per gram dry fermented substrate (U/gds).

**Morphology and cultural characteristics.** The morphology of the spore chain and the spore surface ornamentation of strain NEAE-82 were examined on inorganic salt/starch agar after 14 days at 30°C. The gold-coated dehydrated specimen can be examined at different magnifications with Analytical Scanning Electron Microscope Jeol JSM-6360 LA operating at 20 Kev at the Central Laboratory, City for Scientific Research and Technological Applications, Alexandria, Egypt. Aerial spore-mass color, substrate mycelial pigmentation and the production of diffusible pigments were observed on ISP medium 1–7 as described by Shirling and Gottlieb.

**Physiological characteristics.** Physiological characteristics were performed following the methods of Shirling and Gottlieb. The ability of the organism to inhibit the growth of four bacterial (Pseudomonas aeruginosa, Staphylococcus aureus, Escherichia coli, or Klebsiella), and five fungal strains (Rhizoctonia solani, Alternaria solani, Bipolaris oryzae, Fusarium oxysporum, Fusarium solani) was determined.

**16S rRNA sequencing.** The PCR amplification reaction and sequencing were performed in accordance with the methods described by El-Naggar et al. Sequencing product was deposited in the GenBank database under accession number KJ467538.

**Sequence alignment and phylogenetic analysis.** The partial 16S rRNA gene sequence of strain NEAE-82 was aligned with the corresponding 16S rRNA sequences of the type strains of representative members of the genus Streptomyces retrieved from the GenBank, BLAST, DDBJ and PDB databases by using BLAST program (www.ncbi.nlm.nih.gov/BLAST) and the software package MEGA4 version 2.1 which was used for multiple alignment and phylogenetic analysis. The phylogenetic tree was constructed with the neighbor-joining algorithm based on the 16S rRNA gene sequences of strain NEAE-82 and related organisms.

**Purification of L-asparaginase from Streptomyces fradiae NEAE-82.** All purification steps were carried out at 4°C using crude enzyme extract. The extracellular crude enzyme was prepared at the end of the fermentation period by centrifugation at 11,000 × g for 30 min. The cell free supernatant was used as the crude enzyme preparation. Finely powdered ammonium sulfate was slowly added to the clear supernatant obtained after centrifugation to reach 45% saturation and incubated overnight. The precipitate was collected by centrifugation at 11000 × g for 30 min, while the supernatant was brought to 55–85% saturation with ammonium sulfate. The solution was then centrifuged at 11,000 × g for 30 min. The precipitates were collected separately by centrifugation, dissolved in a minimal amount of 50 mM Tris-HCl buffer pH 8.4 and dialyzed overnight against the same buffer. After dialysis, the samples were used for protein estimation and enzyme assay by the method of Lowry et al. and direct Nesslerization method, respectively and stored at 4°C for further purification.
The dialyzed enzyme solution obtained from the previous step was loaded into a diethylaminoethyl (DEAE) Sepharose CL-6B column (2.3 cm × 12 cm) that was pre-equilibrated with 50 mM Tris–HCl buffer (pH 8.4). The column was washed with two column volumes of the above buffer and the adsorbed protein was isocratically eluted using a 0.5 M NaCl in 50 mM Tris–HCl (pH 8.4). Fractions were collected at a flow rate of 1 ml min⁻¹ (each fraction containing 2 ml) using a fraction collector (Bio-Rad). All chromatographic runs were monitored for protein absorbance at 280 nm. The fractions were collected and examined for enzyme activity and protein content by procedures described elsewhere. Fractions showing high L-asparaginase activity were collected for further use.

Characterization of L-asparaginase enzyme. The effect of the incubation time on L-asparaginase activity was studied by incubating the reaction mixture for different times (10, 20, 30, 40, 50, 60, 70 and 80 min), then the activity of the enzyme was determined. The optimum pH of L-asparaginase activity was studied; L-asparaginase enzyme was pre-incubated with 0.05 M buffers over a range of pH 4–10 under assay conditions, and the amount of ammonia liberated was determined. The buffers used were citric acid-Na2HPO4 (pH 4.5–7.5), Tris-HCl (pH 8.5), and glycine-NaOH (pH 9.5–10.5). The optimum temperature for L-asparaginase activity was determined by incubating the assay mixture at different temperatures ranging from 25 to 60 °C in 0.05 M Tris–HCl buffer under assay conditions. Effect of different substrate concentration on L-asparaginase activity was determined by incubating 0.1 ml from the purified enzyme with different concentration of the specific substrate (0.02, 0.03, 0.04, 0.05, 0.06, 0.07, 0.08, and 1 M) and then the activity of the enzyme was determined under assay conditions.

Determination of kinetic properties ($K_m$ and $V_{max}$). The reaction kinetics of the purified enzyme was determined from Lineweaver-Burk plots with L-asparagine as substrate under defined assay conditions. The Michaelis–Menten constant ($K_m$) and maximal velocity ($V_{max}$) were determined for the enzyme at each of the measured temperatures using the Michaelis–Menten equation:

$$V = \frac{K_m + [S]}{V_{max}[S]}$$

where $V$ is the reaction velocity (a function of enzyme concentration), $[S]$ is the substrate concentration, $K_m$ is the substrate concentration at half-maximal velocity, and $V_{max}$ is the maximal velocity. $V_{max}$ and $K_m$ values were determined using nonlinear regression.

Enzyme thermal stability. Thermal stability of the L-asparaginase was carried by pre-incubating the buffered enzyme prepared in absence of its substrate for different time interval ranging from 0.0 to 90 min at different temperatures (50, 60, 70 and 80 °C). After incubation, the enzyme was cooled then the residual activities were assayed under the standard conditions.

Molecular weight determination and checking of enzyme homogeneity. The molecular weight of the subunit and the purity of enzyme was checked by performing sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE). SDS-PAGE was performed following the method of Laemmli, with separating acrylamide gel 10% (w/v) and stacking gel 5% (w/v) containing 0.1% (w/v) SDS. The gel was stained with coomassie brilliant blue R-250 and de-stained with a solution of methanol, acetic acid and water in the ratio of 4:1:5. Approximate subunit molecular weight of L-asparaginase was determined using standard wide range molecular weight marker (9–178 kDa).

Cell lines. Cancerous cells. Human epithelial colorectal adenocarcinoma cells (caco2 cells), liver hepatocellular carcinoma cells (HepG2 cells) and human laryngeal carcinoma cells (Hep2 cells).

Non-cancerous cells. Human fibroblast cells (FB).

Cytotoxicity and anticancer assay. In order to determine both safety and the anticancer activities of the purified L-asparaginase on both cancerous (CaCo2, HepG2 and Hep2 cells) and non-cancerous cells (Human fibroblasts), neutral red assay protocol was used. Briefly, 100 μl of each of serially diluted purified enzymes in either RPMI or DMEM media were incubated with about 6 × 10⁴ cell/ml of each cell type on 96-well plates. After 48 hours, the cellular cytotoxic effects were quantified using neutral red assay.

Selectivity index (SI). The degree of anticancer selectivity of the tested enzyme was expressed as the previous report with a minor modification. SI = LC50 of pure compound in a normal cell line/LC50 of the same pure compound in cancer cell line, where LC50 is the concentration required to kill 50% of the cell population.

Conclusion
Production of L-asparaginase using different microbial systems has attracted much attention, owing to the cost-effective and eco-friendly nature. The present study revealed an efficient production of an extracellular L-asparaginase throughout various purification steps from isolate Streptomyces fradiae NEAE-82 under submerged fermentation. From the culture filtrate of this strain, L-asparaginase enzyme was purified by column chromatography and molecular weight was determined by SDS-PAGE. The enzyme substrate specificity, kinetic parameters were also determined. Furthermore, high catalytic activity of the enzyme over a wide range of pH and temperature, and its considerable stability, makes it highly favorable for use as a potent anticancer agent, and for other applications in healthcare industry.
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Author Contributions
N.E.E. proposed the research concept, designed the experiments, providing necessary tools for experiments, experimental instructions, conducted most of the experiments, analyzed and interpreted the data and wrote the manuscript. S.F.D. designed and conducted some of the experiments, provided some instruments, experimental instructions, conducted most of the experiments, analyzed and interpreted the data and wrote the manuscript. All authors read and approved the manuscript.

Additional Information
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