Anticancer Properties of N,N-dibenzylasparagine as an Asparagine (Asp) analog, Using Colon Cancer Caco-2 Cell Line

El-Shimaa A. Mohamed¹, Khalid Bassiouny¹, Abeer A. Alshambky¹,², Hany Khalil¹*  

Abstract  

Objectives: This study was conducted to investigate the potential anticancer properties of N,N-dibenzyl asparagine (NNDAsp), as an Asparagine (Asp) analog, using colon cancer Caco-2 cell and the normal NCM-460 cell line. Methods: Cell viability rate and levels of produced lactate dehydrogenase (LDH) were achieved upon treatment with NNDAsp compared to Asp treatment using MTT assay and LDH production kit. The protein expression profile of asparagine synthetase (ASNS) was achieved by using ELISA and flow cytometry assay. The levels of released inflammatory cytokines, including interleukin-1 alpha (IL-1α) and IL-1 beta (IL-β), were monitored using an ELISA assay. Results: Our findings showed significant inhibition of colon cancer cell proliferation accompanied by a high level of produced LDH in a dose-dependent of an NNDAsp treatment without detectable toxic effect in normal cells. Interestingly, NNDAsp showed competitive inhibition of ASNS protein expression, in almost 3% of stained cancer cells, compared to 18% and 35% of untreated cells and cells pre-treated with Asp, respectively. Likewise, the concentration of ASNS protein was dramatically depleted in a dose and time-dependent of NNDAsp treatment in comparison with Asp treatment indicated by ELISA assay. Furthermore, as an apoptotic indicator, the expression of P53 and Caspase 3 (Caps3) was significantly increased in Caco-2 cells treated with NNDAsp at both RNA and protein levels. In contrast, their expression was markedly depleted in Asp-treated cells. In addition, the expression of both IL-1α and IL-1β was markedly increased in Caco-2 cells in a dose and time-dependent of NNDAsp exogenous treatment. Moreover, targeting of ASNS by the Asp analog, NNDAsp, was further confirmed by the docking analysis of inhibitors ligands and crystal structure of ASNS protein. Conclusion: These data provide evidence for the effectiveness of NNDAsp in cancer treatment via selective degradation of ASNS protein expression in colon cancer cells.  

Keywords: Asparagine-N, N-dibenzyl asparagine- asparagine synthetase- Interleukin 1α  

Introduction  

Cell proliferation is one of the most complicated cellular processes that lead to cell division and increasing the copies of each living cell. The mechanisms by which the normal cells can successfully replicate include cyclin-dependent kinases (CDKs), and the epidermal growth factor receptor (EGFR) (Wee and Wang, 2017). The EGFR particularly stimulates downstream targets of the pro-oncoproteins family such as RAS-RAF-MEK-ERK and AKT-PI3K-mTOR signaling pathways (ZHANG and LIU, 2002). In cancer cells, the overexpression of EGFR, and the mutant downstream effectors such as KRAS, BRAF, PIK3CA, P53 and PTEN have been linked to a dynamic change that occurs in cancer cells leading to drug resistance, distinct oncogenicity alteration, and malignancy transformation (Sobani et al., 2016; Balci-Ercin et al., 2020). Colon cancer is one of the most commonly diagnosed cancers and a leading cause of cancer death worldwide. More than one million new cases are recognized yearly with a mortality rate of 60% worldwide, of which cancer metastasis is the leading cause of death (Ferlay et al., 2010; Maher et al., 2020). Indeed, one-third of patients with colon cancer have synchronous or metachronous metastasis. The five-year overall survival rate of patients with primary colon cancer was 80–90%, despite being reduced to 40–60% in patients with advanced nonmetastatic tumors and further decreased to 5–10% in patients with metastatic tumors (Wang et al., 2015). Colorectal cancer (CRC) is the type of cancer that starts in the colon or rectum known as bowel cancer or colon cancer (Kim and Chang 2014). Noteworthy, CRC
begins like a benign tumor, converted to a malignant tumor invading normal tissue and ultimately migrates all over the body. There are many causes of CRC, including obesity, smoking, lifestyle, aging, inflammatory bowel disease, and genetic syndromes (Durko and Malecka-Panas 2014; Simonian et al. 2018).

The asparagine (Asp) is a crucial amino acid in glycopeptide bonds by stimulating N-glycosyl linkage to the sugar rings. L-asparagine (Asp), analog to L-aspartic acid, is one of the nonessential amino acids with a terminal amine group derived from glutamate in the reaction of glutamine to yield Asp in the presence of ATP (Sensei et al., 1991; Yoo et al., 2020). Typically, asparagine synthetase (ASNS) is an enzyme found in all mammalian organs required to convert glutamine to glutamate and aspartate to Asp-dependent ATP molecules (Lomelino et al., 2017). ASNS protein encoded by a gene located on chromosome 7. Notably, the gene of expression ASNS and its protein profile and activity is highly related to cellular stress and drug resistance mechanism of so many diseases and medical disorders. For instance, in acute lymphoblastic leukemia, elevated ASNS protein level in children is associated with the asparaginase-resistant case (Chiu et al., 2020). Several studies also indicated the crosslink between elevated ASNS expression and asparaginase efficacy in treating certain solid tumors (Zhang et al., 2014; Lomelino et al., 2017; Jiang et al., 2021). Like other amino acids, Asp plays a crucial role in cancer development as a building block required for protein synthesis (Jiang et al., 2021). Mechanistically, ASNS and Asp inhibit glutamine-depletion-induced endoplasmic reticulum stress in brain cancer cells (Pavlova et al., 2018). In breast cancer cells, Asp can promote epithelial cell proliferation in the disturbance of exogenous glutamine supply (Zhang et al., 2014). Interestingly, evidence indicated the ability of Asp to enhance the expression of glutamine synthetase (GLUL) resulted in increasing glutamine de novo biosynthesis (Pavlova et al., 2018). Therefore, targeting amino acids such as Asp and amino acid synthetase like ASNS has been proved to be helpful pre-clinical therapy in various types of cancer.

The aim of this study was to investigate the potential anticancer properties of N,N-dibenzyasparagin (NNDAsp), as an Asp analog, in colon cancer cell line in-vitro. The cytotoxicity of NNDAsp in the Caco-2 cell line and the normal colon mucosal epithelial cells, NCM-460 cell line, was compared. The expression profile of ASNS was also assessed in response to NNDAsp and Asp treatment to figure out the potential molecular function of NNDAsp as an anticancer agent.

Materials and Methods

Cell lines
Colon cancer epithelial cell line Caco-2 cells were kindly provided by (VACSERA, Giza, Egypt). Cells were grown in Roswell Park Memorial Institute (RPMI) 1640 media supplemented with 25 mM HEPS, 4 mM L-glutamine, 4 mM sodium pyruvate, and 10% of heat-treated bovine serum albumin (BSA). The normal human colon mucosal epithelial cell line (NCM-460 cells) was grown in RPMI media that contains 4 mM L-glutamine and 10% BSA. All cell lines were incubated at 37°C and a relative humidity of 95% (Khalil et al., 2017; Abd El Maksoud et al., 2019a). The cultured cells’ imaging was determined using inverted microscopy with a Zeiss A-Plan 10X.

Preparation of amino acids stocks
To prepare the Asp and NNDAsp in suitable form, 20 mg of each candidate was dissolved in PBS supplemented with 1% DMSO, and different dilutions has been prepared, including 5, 2.5, 1.25 mg/ml of dissolved buffer. The final extract was incubated at room temperature (R.T.) till used.

Proliferation assay
To achieve cell proliferation, cells were seeded in triplicate in 96-well plates at 10x103 cells per well or in duplicate in a 6-well plate at 10X104 cells per well. Cell morphology was assessed using an inverted microscope. The number of living cells was accounted by using a hemocytometer. Accordingly, the old media was removed, and then the cells were washed twice by PBS and trypsinized by adding an appropriate volume of trypsin followed by 3 min incubation at 37°C. Finally, an appropriate volume of complete RPMI media was added to the trypsinsized cells, and the number of cells was accounted using the inverted microscope (Tahe et al., 2021).

Cytotoxic concentration 50% (CC50)
The Asp and NNDAsp were tested for their anticancer properties and calculated the potential CC50 in Caco-2 and NCM-460 cells. Therefore, the cells were cultured in triplicate in 96-well plates in a density of 10X103 cells/well and were incubated overnight at 37°C and humidity conditions. The cells were treated with different concentrations of each component (20-1.25 mg/ml) followed by overnight incubation. The cell viability rate and cytotoxic concentration were monitored by using MTT colorimetric assay kit (Sigma-Aldrich, Germany). Accordingly, the old media was removed from treated cells seeded in 96-well, and PBS washed the cells, and then 100 µl complete RPMI media was added to each well. Subsequently, 10 µl MTT solution was added to each well, and the plate was incubated for 2 hours at 37°C. Finally, 100 µl SDS-HCl solution was added to each well in the plate which then was incubated for 4 hrs at 37°C. Cell viability was then assessed based on the amount of converted water-soluble MTT to an insoluble formazan which is then solubilized and determined by optical density at 570 nm (El-Fadl et al., 2021).

Lactate dehydrogenase (LDH) production
LDH assay kit (Abc-65393) was used to assess LDH production in the fluid media collected from cultured-treated cells. According to the manufacturing procedures, 100 µl of lysed cells was incubated with 100 µl LDH reaction mix for 30 min at room temperature. A plate reader was used to quantify LDH activity at OD 450 nm.
Flow cytometry analysis

Flow cytometry was used to evaluate the protein expression profile of phosphorylated P53 (Pho-P53), Casp3 and ASNS enzyme in treated Caco-2 cells. Accordingly, drug-treated cells were washed by phosphate buffer saline (PBS) and were trypsinized for 3 min. The complete RPMI medium was added to the trypsinized cells, which were then centrifuged for 3 min at 1500 rpm. The supernatant was discarded, and the pellet was resuspended in PBS for washing and resuspended in PBS with 0.2% formaldehyde for fixation. The cells were resuspended in PBS for permeabilization, including triton-x-100 (0.1%), and incubated for 3 min. For staining of Pho-P53, the cells were resuspended and incubated for 2 hrs at R.T. in the PBS supplemented with 1% BSA and the diluted mouse monoclonal anti-pho-P53 (Sigma-Aldrich, Germany). After washing, the cells were centrifuged and resuspended in the PBS that contains 1% BSA and 1-1000 secondary antibody donkey anti-mouse IgG (Alexa Fluor 594, Invitrogen, Germany). The same conditions were followed in staining of p-ho-Casp3 protein in treated cells using rabbit polyclonal anti- p-ho-Casp3 (PromoCell, Germany) and goat anti-rabbit IgG (Alexa Fluor 488, Abcam, USA). For staining ASNS enzyme, the rabbit monoclonal primary antibody (mAb #92479, Cell Signaling Technology, USA) and goat anti-rabbit IgG (Alexa Fluor 488, Abcam, USA) were used. Finally, the flow cytometry assay (BD Accuri 6 Plus) was used to assess the protein levels using a resuspended pellet in 500 μL PBS (Abd El Maksoud et al., 2019b; Hamouda et al., 2021).

Enzyme-linked immunosorbent assay (ELISA)

Human asparagine synthetase (ASNS) ELISA Kit (Abbexa, Cambridge, UK) has been used. The microplate provided in this kit has been pre-coated with an antibody specific to ASNS. Standards or samples were then added to the appropriate microplate wells with a biotin-conjugated antibody specific to ASNS. Next, Avidin conjugated to Horseradish Peroxidase (HRP) is added to each microplate well and incubated. After TMB substrate solution was added, only those wells containing ASNS, biotin-conjugated antibody, and enzyme-conjugated Avidin will exhibit color changes. The enzyme-substrate reaction was terminated by adding sulphuric acid solution, and the color change was measured spectrophotometrically at a wavelength of 450 nm +/- 10nm. The concentration of ASNS in the samples is then determined by comparing the O.D. of the samples to the standard curve. ELISA assay was also used to quantify the released interleukin-1α (IL-α) and IL-1β using human ELISA kits (Abcam 46028 and Abcam 46052, respectively). According to the manufacture protocol, cells cultured in 96-well plates were overnight incubated. Then the cells were treated with different concentrations of each amino acid followed by an incubation period of (0, 6, 12, 24, 36, 48, and 72 hrs). At each time point, the cells were lysed using 1X cell lysis buffer (Invitrogen, USA). Then, 100 μl of the lysed cells were transferred into the ELISA plate reader and incubated for 2 hrs R.T. with 100 μl control solution and 50μl 1X biotinylated antibody. Then 100 μl of 1X streptavidin-HRP solution was added to each well of samples and incubated for 30 min in the dark. 100 μl of the chromogen TMB substrate solution was added to each well of samples and incubated for 15 min at R.T., away from the light. Finally, 100 μl stop solution was added to each well of samples to stop the reaction. The absorbance of each well was measured at 450 nm (Khalil H et al., 2017; Khalil et al., 2020a).

Quantitative real-time PCR (qRT-PCR)

The quantification analysis of gene expression was detected using qRT-PCR. The total cellular RNA was obtained using TriZol (Invitrogen, USA) and purified using an RNA purification kit (Invitrogen, USA). Complementary DNA (cDNA) was synthesized from 1 μg of total RNA using M-MLV reverse transcriptase (Promega, USA). The P53 and Caspase 3 (Casp3) mRNA expression quantification analyses were achieved using QuantiTect-SYBR-Green PCR Kit (Qiagen, USA) and the specific primers listed in Table 1. The housekeeping glyceraldehyde 3-phosphate dehydrogenase (GAPDH) expression level was used for normalization in the real-time PCR data analysis. The PCR reaction system contained 10 μl SYBR green, 0.25 μl RNase inhibitor (25 U/µl), 0.2 μM of each primer, 2 μL of synthesized cDNA, and nuclease-free water up to a final volume of 25 μL. The following PCR conditions were used; 94°C for 5 min, 35 cycles (94°C for 30 sec, 60°C for 15 sec, 72°C for 30 sec) (Khalil et al., 2019; El-Fadl et al., 2021).

Data analysis

Microsoft Excel was used to prepare all histograms and charts. Delta-Delta Ct analysis was used in the quantification analysis of mRNA delivered from qRT-PCR assay based on the following equations: (1) $\Delta \Delta Ct = Ct$ value for gene - Ct value for GAPDH, (2) $\Delta (\Delta \Delta Ct) = \Delta Ct$ value for experimental - $\Delta Ct$ for control, (3) Quantification fold change = (2 - delta-delta ct ) (Rao et al., 2013; Khalil et al., 2017). The student’s two-tailed t-test was used for statistical analysis. P-value ≤ 0.05 was considered statistically significant.

Results

NNDAsp disturbs Caco-2 cell proliferation without affecting NCM-460 cell proliferation

To test the cytotoxic effect of NNDAsp cell viability rate, MTT assay and relative produced LDH were achieved in Caco-2 and NCM-460 that pretreated with different concentrations of either Asp or NNDAsp. As shown in Figure 1A and Table 2, the mean absorbance values revealed cell viability rate of Caco-2 cells was markedly decreased in a dose-dependent manner of NNDAsp, while showed increasing rat in response to Asp treatment. Interestingly, in both cancer and normal cells, the mean absorbance of treated cells showed constant values in
response to Asp and NNDAsp (Figure 1B and Table 2). These findings indicate that NNDAsp can disturb cancer cell proliferation mainly with a cytotoxic concentration of 50% (CC50) with 4 mg/ml. Further, the relative LDH production was significantly increased in cancer cells upon treatment with 4 mg/ml of NNDAsp for 24 hours compared with untreated cells and cells treated with Triton X-100, as a toxic compound, without detectable cytotoxic effect in normal cells (Figure 1C, Table 3, and Table 4). Together, these data suggest the potential anticancer properties of NNDAsp and further confirm the crucial role of Asp in cell proliferation.

**NNDAsp affects cancer cell morphology and the number of survived cells**

Cell morphology and the number of survived cells were assessed in both treated cells with Asp and NNDAsp using an inverted microscope. Markedly, the cell morphology revealed a disturbing proliferation of Caco-2 cells treated with 4 mg/ml NNDAsp compared with the cell morphology of other treated cells (Figure 2A). The number of surviving cells significantly decreased in response to NNDAsp treatment only in colon cancer cells. At the same time, the same concentration of Asp

### Table 1. Oligonucleotides sequences used for mRNA Quantification of Indicated Genes

| Description   | Primer sequences 5’-3’ |
|---------------|------------------------|
| Casp3-forward | GGACACGAGTTACAAAATGGATTA |
| Casp3-reverse | GGCAGGGCCTGAATGTAAGGAG |
| P53- forward  | GCGAGCCTGCCAACAACACA   |
| P53- reverse  | GTGCACGGTCTTGGTGCCTT   |
| GAPDH-forward | TGCCATTGTGGGAAGGCTCTA  |
| GAPDH-reverse | TGATGCAGGGATGATGTTCT   |

### Table 2. Cell Viability Rate of Treated Caco-2 Cells

| Treatment | Concentration (mg/ml) | Mean Absorbance (570nm) | P-values |
|-----------|-----------------------|-------------------------|----------|
|           |                       | NCM-460 cells | Caco-2 cells | NCM-460 cells | Caco-2 cells |
| Asp       | 0.0                   | 0.57±0.1 | 0.57±0.1 | > 0.05 | > 0.05 |
|           | 1.25                  | 0.70±0.06 | 0.70±0.06 | > 0.05 | > 0.05 |
|           | 2.5                   | 0.80±0.11 | 0.84±0.11 | < 0.05* | < 0.05* |
|           | 5                     | 0.74±0.1 | 0.89±0.08 | < 0.05* | < 0.01** |
|           | 10                    | 0.86±0.08 | 0.89±0.06 | < 0.01** | < 0.01** |
|           | 20                    | 0.89±0.07 | 1.01±0.02 | < 0.01** | < 0.01** |
| NNDAsp    | 0.0                   | 0.65±0.13 | 0.6±0.13  | > 0.05 | > 0.05 |
|           | 1.25                  | 0.58±0.1 | 0.60±0.13 | > 0.05 | > 0.05 |
|           | 2.5                   | 0.5±0.1  | 0.48±0.13 | > 0.05 | > 0.05 |
|           | 5                     | 0.54±0.08 | 0.23±0.1 | > 0.05 | < 0.01** |
|           | 10                    | 0.64±0.13 | 0.12±0.1 | > 0.05 | < 0.01** |
|           | 20                    | 0.74±0.13 | 0.08±0.02 | > 0.05 | < 0.01** |

¥, DMSO; *, Indicates significant P values ≤ 0.05; **, Indicates high significant P values ≤0.01

![Figure 1](image1.png)

**Figure 1.** Cell Viability and Cytotoxic Effect of Asp and NNDAsp Treatment. (A) Cell viability rate of Caco-2 cells that were pretreated with indicated concentrations of either Asp or NNDAsp for 24 hrs compared with DMSO (0 concentration). (B) Cell viability rate of NCM-460 cells that were pretreated with the same concentration of Asp or NNDAsp for 24 hrs compared with DMSO (0 concentration) using MTT assay. (C) Relative LDH production from treated cells compared to Triton 100-X treated cells and untreated cells (N.T.). Error bars indicate the S.D. of four different replicates. The student's t-test was used to determine the significance of differentiated values. (*) indicates P-values ≤0.05, and (**) indicates the P ≤0.01.

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showed an increasing number of survived cells (Figure 2B and Table 5). These findings indicate the severe effect of NNDAsp treatment in the Caco-2 cells. In contrast, Asp treatment stimulates cell proliferation without any detectable toxic effect.

Figure 3. Expression Profile of P53 and Casp3 as Apoptosis Regulators. (A) Quantification of the mRNA expression level of the P53 gene indicated by fold change in treated Caco-2 cells compared with untreated cells (N.T.) and DMSO-treated cells. (B) The mRNA expression level of Caps3 gene was quantified as indicated by fold change in treated Caco-2 cells compared with untreated cells (N.T.) and DMSO-treated cells. Error bars indicate the S.D. of two independent experiments. The student's two tails t-test was used to determine the significance of differentiated Ct values. (*) indicates P-values ≤0.05, and (**) indicates the P ≤0.01.
NNDAsp stimulates programmed cell death of treated colon cancer cells

To confirm the cooperation of NNDAsp in cancer cell death, the expression profile of P53, as a tumor suppressor gene, and Casp3, as an indicator for apoptotic signaling, was monitored in treated Caco-2 cells at both RNA and protein levels. Expectedly, the relative expression of both P53 and Casp3 was dramatically unregulated, at the RNA level, in cells treated with NNDAsp compared to control cells and cells treated with Asp (Figure 3A and B). The statistical analysis presented in Table 7 indicated that the upregulation of P53 and Casp3 gene
expression was highly significant since P values were equal to 0.002 and 0.01, respectively. Furthermore, flow 
cytometry was used for quantification of phos-P53 and 
Casp-3 proteins at a single-cell level via a combination 
of specific primary antibodies to stain treated cells. To 
quantify the corresponding proteins profile in treated cells, 
the percentage of positive events was correlated to the 
relative fluorescent intensity of the Alexa Fluor signal of 
secondary antibodies. Interestingly, the kinetic expression 
profile of both phos-P53 and Casp3 proteins was markedly 
increased more up to 20%, while their kinetic expression 
was strongly decreased less than 5% of stained-control 
cells and cells treated with Asp (Figure 3C). Together, 
these data suggest that NNDAsp can restore the expression 
of P53 and stimulate the expression of Casp3 TET3 in 
both RNA and protein levels to produce programmed cells 
death in treated cells (PCD) in infected cells.

NNDAsp regulates the expression profile of ASNS enzyme 
in a time and dose-dependent manner

To address the connection of amino acids treatment 
and the expression profile of ASNS protein in treated cells, 
ASNS protein expression was measured in different time 
points and various concentrations upon treatment using 
ELISA assay. Compared with Asp, NNDAsp treatment 
showed a potent reduction of ANSN protein concentration 
in a time and dose-dependent manner (Figure 4A and B). Furthermore, the quantification analysis of ASNS protein

Table 4. LDH Production in Treated Normal Cells (NCM-460 cells)

| Treatment | NCM-460 cells | Caco-2 cells | NCM-460 cells | Caco-2 cells | NCM-460 cells | Caco-2 cells |
|-----------|---------------|--------------|---------------|--------------|---------------|--------------|
| Mean absorbance | 0.06±0.03 | 0.27±0.03 | 0.05±0.01 | 0.06±0.02 |
| Relative LDH production | 1 | 1.6 | 6.5 | 1.2 | 1.44 |
| P values | 0.14 | 0.001** | 0.37 | 0.77 |

Table 5. Number of Survived Cells Upon Treatment

| Treatment | NCM-460 cells | Caco-2 cells | NCM-460 cells | Caco-2 cells | NCM-460 cells | Caco-2 cells |
|-----------|---------------|--------------|---------------|--------------|---------------|--------------|
| Mean absorbance | 0.04±0.01 | 0.06±0.03 | 0.27±0.03 | 0.05±0.01 | 0.06±0.02 |
| P values | 0.14 | 0.001** | 0.37 | 0.77 |

Table 6. Quantification Analysis of P53, and Casp3 Gene 
Expression in Caco-2-Treated Cells

| Genes | Treatment | Expression fold changes | P-values |
|-------|-----------|-------------------------|----------|
| P53   | NT        | 1.00±0.00               | > 0.05   |
|       | DMSO      | 0.8±0.17                | > 0.05   |
|       | Asp       | 0.18±0.07               | > 0.05   |
|       | NNDAsp    | 8.6±0.26                | < 0.01** |
| Casp3 | NT        | 1.00±0.00               | > 0.05   |
|       | DMSO      | 1.04±0.5                | > 0.05   |
|       | Asp       | 0.42±0.04               | > 0.05   |
|       | NNDAsp    | 2.37±0.03               | < 0.05*  |

Figure 6. Binding Discrimination of Amino Acid Ligands with ASNS Protein. (A and B) The pharmacophore mapping 
snapshot indicates the possible hydrophobic interaction and binding affinity between Asp, NNDAsp, respectively, and 
the ASNS chain protein crystal structure.
expression using flow cytometric assay showed a potent reduction of ASNS protein expression, less than 5%, in stained Caco-2 cells treated with 4 mg/ml NNDAsp. While showed more than 18% in control cells, 19% in DMSO-treated cells, and 35% in cells treated with the same concentration of Asp (Figure 4C). Collectively, these findings proved the targeting of ASNS protein by NNDAsp in a time and dose-dependent manner in the treated Caco-2 cell line.

**NNDAsp stimulates IL-1α and IL-1β expression in colon cancer cells**

To determine the correlation between NNDAsp treatment and produced inflammatory cytokines from treated cells, IL-1α and IL-1β were measured in a time-course experiment using ELISA assay. Interestingly, NNDAsp treatment showed a high production level of both IL-1α and IL-1β in a time-dependent manner. In contrast, the production level of both interleukins was dramatically reduced in treated cells with Asp and control-treated cells (Figure 5A and B). Together, these data further confirm the impact of NNDAsp in regulating the production of proinflammatory cytokines from treated cells and reveal its ability to stimulate PCD in colon cancer cells.

**Binding Based-Cross Docking Analysis of Asp and NNDAsp Ligands and ASNS crystal structure**

The docking analysis presented in Figure 6 reveals that the ASNS protein residues ARG 184 and ARG 203 formed a hydrogen bond with an Asp ligand molecule (A). Meanwhile, PHE 352 and ASP 86 in ASNS protein formed hydrogen bonds with the NNDAsp molecule (B). These results indicate the possible interaction and direct interference of NNDAsp and Asp with ASNS protein in treated cells.

**Discussion**

The role of ASNS and amino acids in hallmarks of tumor malignancy has been reported in several studies (Kuo et al., 2021; Wei et al., 2021). In the current work, we investigated the molecular influence of NNDAsp, as a modified amino acid, in the proliferation attitude of colon cancer Caco-2 cells compared to normal human colon mucosal epithelial cells (NCM-460 cells). NNDAsp demonstrated the ability to induce PCD in colon cancer cells via rescue P53 and Caps3 gene expression in treated cells. Significantly, NNDAsp reduced the expression profile of the ASNS enzyme, which is responsible for the generation of aspartate and glutamate and contributes to the drug resistance mechanism during cancer therapy. Our findings provide evidence for the potential anticancer properties of NNDAsp with minimal cytotoxic outcomes in normal cells.

During the imbalanced availability of amino acids, the ASNS gene is targeted by two signaling pathways to ensure cell survival: the Unfolded Protein Response (UPR) (Kilberg et al., 2009) and the Amino Acid Response (AAR) (Kilberg et al., 2012). Under the endoplasmic reticulum (ER) stress and through the activation of the PKR-like ER (PER) and GCN2 protein kinases, both UPR and AAR pathways converge on the phosphorylation of the α-subunit of the initiation factor eIF2. Subsequently, the eIF2 effector provokes the depletion of global protein synthesis and the translation of selected mRNAs, including the ATF4 transcription factor. Noteworthy, ATF4 is a crucial trans-activator for ASNS induction by binding to an enhancer element within the ASNS promoter (Balasubramanian et al., 2013). Recent evidence demonstrated the translational reprogramming in ASNS-depleted cancer cells that depends on the increase of the MAPK-interacting kinase 1 (MNK1) and eukaryotic translation initiation factor 4E (eIF4E), resulting in increasing ATF4 translation and ASNS expression (Pathria et al., 2019). Additionally, ASNS transcription is also affected negatively by other factors like P53, a tumor suppressor gene (Chiu et al., 2020). Likewise, we hypothesized that NNDAsp treatment could negatively regulate ASNS protein expression in colon cancer cells via direct interaction with its synthetase sites. As a negative correlation, the depletion of ASNS by NNDAsp in colon cancer cells activated the expression of P53 and stimulated the programmed cell death in treated cells. The human ASNS enzyme includes the C-terminal domain, similar to ASNS in other organisms, and the N-terminal glutaminase domain. The glutaminase domain has a topology like other amidotransferases. In addition, the amino acids in the synthetase site are, for the most part, conserved in human and bacterial ASNS. Also, other preserved amino acid residues are recognized in the interface of ASNS domains (Zhu et al., 2019). Indeed, we identified two binding sites at the N-terminal domain of the ASNS enzyme by which NNDAsp binds with amio acide residues; phenylalanine and asparagine indicated by docking analysis. Accumulating evidence showed the association between Raf/MEK/ERK signaling and the expression of Asp biosynthesis enzyme ASNS, demonstrating the sensitivity of melanoma and pancreatic cancer cells to Asp depletion, resulting in disturbing cancer cell proliferation (Pathria et al., 2019). Also, ASNS depletion has been reported to inhibit the panel of breast cancer, prostate cancer, and melanoma cell proliferation (Pathria et al., 2019). Notably, the mitogen-activated protein kinase (MAPK), including Raf/MEK/ERK pathway, serves as a critical signal transducer of receptor tyrosine kinases. The Ser/Thr kinase Raf protein (c-Raf-1, Raf-B, or Raf-A) activates the downstream kinase MEK1 and MEK2, which, in turn, phosphorylate Ser/Thr kinases ERK1 and its homolog ERK2 (Shaul and Seger, 2007; Eliman et al., 2020). The Raf/MEK/ERK pathway regulates cell survival, cell cycle progression, and differentiation. Furthermore, the Raf/MEK/ERK pathway can also modulate the activity of many proteins involved in the apoptotic signal, such as Bim, Bax, Casp-9, Casp-3, and P53, which, in turn, regulate cellular autophagy and result in PCD (McCubrey et al., 2007; Khalil et al., 2020b; El-Fadl et al., 2021).

Collectively, our findings further proved ASNS as an attractive candidate to prevent cancer proliferation and stimulate the expression of the tumor suppressor gene, P53, which in turn activates apoptotic signaling in colon cancer cells.
In conclusion, the expression of ASNS in cancer cells is increased to fill the cell need for Asp, which cooperates in many biological processes related to cell proliferation and cancer development. Thus targeting Asp by asparaginase is affected in cancer therapy; however, ASNS expression was linked to the asparaginase-resistant mechanism during therapy. Therefore, this study demonstrates that NNDAsp, an Asp analog, can disrupt colon cancer cell proliferation without affecting the normal human colon mucosal epithelial cell line. NNDAsp successfully induced PCD in cancer-treated cells by restoring P53 gene expression and regulating apoptotic signaling in treated cells. Importantly, our findings proved that NNDAsp targets ASNS enzyme indicated by ELISA assay and flow cytometric analysis in the Caco-2 cell line.

Docking analysis confirmed the possible interaction and binding affinity between NNDAsp and ASNS protein domain structure. Furthermore, NNDAsp stimulates the production level of proinflammatory cytokines IL-1α and IL-1β that facilitate the PCD in cancer-treated cells. Together, these data suggested that NNDAsp is an attractive agent that can regulate colon cancer cell proliferation without detectable toxic effects in normal cells and indicating the potential role of NNDAsp indirectly to overcome the asparaginase-resistant mechanism during cancer therapy.

Author Contribution Statement

El-Shimaa Salah and Abeer Alshambky provided the experiments and helped in writing the manuscript. Khalid Bassiony helped in supervision and data analysis. Hany Khalil designed the research plan, supervised overall research, provided and interpreted data and wrote the manuscript.

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Ethical statement
This study was approved by the ethical committee of Genetic Engineering and Biotechnology Research Institute, University of Sadat City, Egypt.

Availability of data and materials
The data supporting this study’s findings are available from the corresponding author upon reasonable request.

Conflicts of interest
All authors declare that there are no conflicts of interest.

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