IN VITRO ANTI Can cer A CTIVITY OF NANA ZO XD DRUG AGA IN TS COLORECTAL CANCER CELL LINE AND ITS MOLECULAR PATHWAYS


1- MSc student at Department of Biotechnology, Faculty of Science, Cairo University, Cairo, Egypt.  
2- Drug Bioassay-Cell Culture Laboratory, Pharmacognosy Department, National Research Center, Dokki, Giza, 12622, Egypt.  
3- Department of Zoology, Faculty of Science, Cairo University, Cairo, Egypt.  
4- Department of Chemistry, Faculty of Science, Cairo University, Cairo, Egypt.  
*Corresponding author’s E-mail: Hallouty68@gmail.com

Submitted August 3, 2021; Accepted October 11, 2021; Published November 21, 2021

SUMMARY

Drug repurposing is gaining popularity in treatment discovery because it is a smart strategy to quickly exploit new molecular targets of a known clinically authorized drug in a variety of diseases, both existing and novel. Some early studies showed that hydroxychloroquine which has long been used for malaria, is used to reduce fever and inflammation of COVID-19 patients. Cancer is a major health issue worldwide. During the past years, some drugs are proved to be anticancer drugs as Raloxifene to treat breast cancer and Aspirin to treat colorectal cancer.

According to the national cancer institute the second most killer cancer is the cancer of colon and rectum around the world. Colorectal cancer is Egypt's seventh most prevalent cancer, accounting for 3.47 % cancers. Colon cancer patients (excluding rectal cancer) were expected to number somewhat more than 3,000. Our aim in this project is to repurpose clinically used antibiotics in the Egyptian market for colorectal cancer. Anticancer efficacy of numerous antibiotic drugs was being tested by screening them in monolayer (2D).

Results showed that Nanazoxid as a promising drug as anti-cancer drug especially for colorectal cancer with IC50 (0.20 μg/ml) and was tested for its selectivity index on human cell line and was subjected to molecular analysis. As a result, an antibiotic drug which is Nanazoxid proved that it could be repurposed as an anti-colorectal therapy with proper further research and development.

Keywords: Colorectal Cancer- Drug Repurposing- -Antibiotics- human cell line-Aspirin- COVID-19-Nanazoxid

INTRODUCTION

The process of discovering new biological targets for existing drugs that have already been approved for the treatment of other diseases or whose targets have already been discovered is known as drug repurposing. This method is not time-consuming, costly, or financially risky. (Padhy and Gupta. 2011) Also, it is able to be relevant on recognized and uncommon illnesses as Covid-19 that's A novel coronavirus (CoV) named ‘COVID-19’ with the aid of using the World Health Organization (WHO) is in price of the modern outbreak of pneumonia that commenced at the start of December 2019 close to in Wuhan City, Hubei Province, China (Hui et al.,2020 ) and the time is so restricted to discover a new technique to deal with COVID-19 however drug repurposing that's the quicker manner helped to locate powerful healing dealers because the antiviral and antimalarial sremdesivir, favipiravir, ribavirin, lopinavir-ritonavir combination, arbidol, tocilizumab, chloroquine and hydroxychloroquine(Singh et al.,2020).

Cancer is a main cause loss of life and morbidity across the world. According to current World Health Organization estimates every year most cancers prevalence in Sub-Saharan Africa is 551 200, with 421 000 deaths in line with year (2008) Low- and middle-profits international locations accounted for nearly 70% of all cancers fatalities (El-Hallouty et al.,2015)

Colorectal cancer is the world's third most frequent cancer and the fourth leading cause of death, accounting for more than 9% of all cancer cases. It has equal risk in both sexes where males have a slightly higher risk for rectal cancer than females. Furthermore, colorectal cancer affects roughly 40/100,000 people in the United States, Australia, and Western Europe, compared to approximately5/100,000 in Africa and portions of Asia. (Haggar and Boushey, 2009).

Nanazoxid drug (nitazoxanide) is the treatment for Cryptosporidium infections (Gargala ,2008) In Africa and Asia, Cryptosporidium infection is the second cause of severe diarrhea in young children (Kotloff et al.,2013) and in this study we discovered the high cytotoxic effect of nanazoxid on colorectal cancer cell line HCT-116 with IC50 0.20 μg/ml and selectivity index 125.41.

Issued by The Egyptian Arab Foundation for Investment, Innovation and Industrial Development (EAFID)
Nanazoxid possess antiproliferative activity according to the flowcytometry results as the cell growth arrest at G1/S phase and induces apoptosis of HCT116 cells by upregulating Bax - P53 - Casp3 genes and downregulating the BCL-2 gene.

MATERIALS AND METHODS

Antibiotics:
Table 1. The antibiotic drugs used in the research were obtained from the Egyptian market

<table>
<thead>
<tr>
<th>Drug</th>
<th>Company</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nanazoxid</td>
<td>Utopia pharmaceutical</td>
</tr>
<tr>
<td>Benzabiotic</td>
<td>Sedico pharmaceutical</td>
</tr>
<tr>
<td>Quinabiotic</td>
<td>Utopia pharmaceutical</td>
</tr>
<tr>
<td>Floxamo</td>
<td>amoun pharm.co</td>
</tr>
<tr>
<td>Tavanic</td>
<td>sanofiwinthropindustrie</td>
</tr>
<tr>
<td>Respenzo</td>
<td>hochster</td>
</tr>
<tr>
<td>Augmentin</td>
<td>GlaxoWellcome</td>
</tr>
<tr>
<td>Rovac</td>
<td>delta</td>
</tr>
<tr>
<td>Ceporex</td>
<td>GLAXO</td>
</tr>
<tr>
<td>Omnicef</td>
<td>Al-Hikma Pharmaceuticals</td>
</tr>
</tbody>
</table>

Cancer cell lines:
In this study, the human colon carcinoma cell line (HCT-116), and a skin normal human cell line (BJ-1) immortalized normal foreskin fibroblast cell line. These cell lines were obtained from Karolinska Institute, Department of Oncology and Pathology (Stockholm, Sweden).

Molecular analysis:
RN easy extraction kit (Qiage, Hilden, Germany), Super Script™ II Reverse Transcriptase (Biorad, USA), iScript One-Step RT-PCR Kit with SYBR® Green (Biorad, USA), Forward and reverse primers of target genes (Sigma, USA). Annexin V-FITC Apoptosis Detection kit: (BioVision, USA)

Cell culture:
HCT116 human colon carcinoma cell line and skin normal human cell line (BJ-1) were maintained in DMEM: F12 Medium/10% FBS. Both cell lines were incubated at 37°C in 5% CO2 and 95% humidity. Cells were sub-cultured using trypsin versene 0.15%.

MTT assay:
After 24hrs of seeding 10,000 HCT-116 cells per well and 50,000 BJ-1 cells per well (in 96-well plates), a100-ppm final concentration of the tested drugs were added in triplicates. The cells were treated for 120hrs. 1 µM Doxorubicin was used as positive control and 0.5% DMSO was used as a negative control. Cytotoxicity was determined using MTT [3-(4, 5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] assay as described by Mosmann in 1983 (Mosmann, 1983). Cytotoxicity was calculated according to the following equation: [1-(av(x))/(av (NC))] *100. Where: Av: average, X: absorbance of sample, NC: absorbance of negative control. Absorbance was measured at 595nm with reference 690nm.

Determination of IC_{50} values:
In case of highly active drugs possessing ≥ 70 % cytotoxicity on colorectal cancer cell line. Different concentrations were prepared for dose response studies. The results were used to calculate the IC_{50} values of each drug using probit analysis and utilizing the SPSS computer program (SPSS for windows, statistical analysis software package / version 9 / 1989 SPSS Inc., Chicago, USA).

Selectivity Index (SI):
The selectivity index (SI) indicates the cytotoxic selectivity (i.e. safety) of the drug against colorectal cancer cell versus normal cells (BJ-1, skin human normal immortalized cell line). SI = IC_{50} of promising drug in a normal cell line/ IC_{50} of the same promising drug in cancer cell line. (Prayonget al., 2008)

B. RNA isolation:
The effect of the drug (Nanazoxid) on the mRNA expression of the Bcl-2, P53, Caspase-3 and Bax was determined by a reverse transcription-polymerase chain reaction (RT-PCR) technique. After the treatment (48 h) [12] of the HCT116 cells with the extract, RNA was extracted from the cells using the RNeasy extraction kit (Qiagen, Hilden, Germany) following the manufacturer’s protocol. Concisely, up to 1 × 106 of the cultured cells were trypsinized, washed with PBS, and collected in an RNase-free centrifuge tube. The sample was then centrifuged at 300 xg for 5 min, and the supernatant was aspirated off. The pellet was disrupted and homogenized in RLT buffer (buffer RLT...
is a lysis buffer for lysing cells and tissues before RNA isolation and simultaneous RNA/DNA/protein isolation.

The homogenized lysate was then transferred into a 2 ml collection tube, and 1 ml of 70% ethanol was added to the lysate and mixed with pipetting. Then, 700 μl of the sample was transferred into an RNeasy spin column placed in a 2 ml collection tube and centrifuged at 10,000 rpm for 15 sec. The flow-through was discarded and 700 μl of the RW1 buffer (buffer RW1 is a proprietary component of RNeasy kits. Buffer RW1 contains a guanidine salt, as well as ethanol, and is used as a stringent washing buffer that efficiently removes biomolecules such as carbohydrates, proteins, and fatty acids) was added to the spin column and centrifuged at 10,000 rpm for 15 s.

The previous step was repeated 2 times using 500 μl of the RPE buffer (buffer RPE is a mild washing buffer and a proprietary component of RNeasy kits. Its main function is to remove traces of salts, which are still on the column due to buffers used earlier in the protocol) instead of the RW1 buffer to wash the spin column membrane. The collection tube was discarded, and a new 1.5 ml tube was used instead, 40 μl of RNase-free water was added to the center of the membrane and centrifuged at 10,000 rpm for 1 min. The spin column was discarded as the collection tube contains the eluted RNA.

C. Gene expression analysis:
After the RNA extraction, one-step RT-qPCR was performed using iScript™ One-Step RT-PCR kit with SYBR® Green (Bio-Rad Inc., CA, USA) which was added to the Master Mix which contained 25 μl 2x SYBR Green®, 1.5 μl forward primer (10 μM), 1.5 μl reverse primer (10 μM), 11 μl nuclease-free water, and 1 μl 50 iScript Reverse Transcriptase in a final reaction volume of 50 μl. Two control reactions were created; one lacking the RNA template and the other lacking the reverse transcriptase enzyme.

The one-step RT-qPCR reaction included an incubation step with the complete reaction mix, iScript Reverse Transcriptase enzyme inactivation step, qPCR step (denaturation, annealing, and extension), and dissociation step. The amplification reactions were carried out using the Rotor-Gene Q Real-Time PCR system (Qiagen, Hilden, Germany).

The primers’ sequences used to quantify the gene expression of the of Bcl-2, P53, Caspase-3, Bax and β-Actin as housekeeping genes are:

- Caspase-3 gene is F: 5’-TGTGGTGTGCTCTGAGCC-3’ and R: 5’-CACGCCATGTCATCAAC-3’.
- Bax gene is F: 5’-ATGTTTTTCTGAGGGCAACTTC-3’ and R: 5’-AGTCCAATGTCGCCCAT-3’.
- β-Actin gene is F: 5’-GTGACATCCACACCCAGAGG-3’ and R: 5’-ACAGGATGTCAAAACTGCC-3’.
- Bcl-2 gene is F: 5’-ATGTTGTTGGGAGACCCTCAAA-3’ and R: 5’GCCGTACAGTTCCACAAAGG-3’.
- P53 gene is F: 5’-ATGTTTTGCAAACCTGGAACG-3’ and R: 5’TGAGCAGCGCTCATGGT-3’.

D. Flow cytometry:
For the measurement of the cellular DNA content, flow cytometric analysis was carried out using the Annexin V-FITC Apoptosis Detection kit (BioVision, CA, USA). After 48 h of treatment, HCT116 cells were trypsinized and washed with serum-containing media. The solution was then centrifuged at 300xg for 10 min, the supernatant was decanted, and the pellet was resuspended in 500 μl of ×1 binding buffer. Then, 5 μl of Annexin V-FITC and 5 μl of propidium iodide were incubated with the cells for 5 min in the dark. The sample was quantified by BD FACSCalibur (BD Bioscience, CA, USA)

Results
A. Cytotoxicity bioassay on HCT116 monolayers:
Data were pooled from the MTT assays where 10 antibiotic drugs were tested on HCT116 lines (100 ppm) the results showed after screening of antibiotics, all are below 70% for the cytotoxicity effect. Only one drug is promising which is Nanazoxid has more than 70% cytotoxicity effect. So it’s the only drug was prepared in different concentration ranges from 100 ppm to 0.048 ppm then the determination of IC50 in comparison with normal cell line (BJ-1)
Figure 1: The results of Nanazoid, Quinabiotic, Benzabiotic, Floxano, Tavenic, Repenzo, Augmentin, Rovac, Ceporex, and Omnicer mean cytotoxicity effect on HCT-116 cells (100 ppm), as shown in the schematic.

Figure 2: The mean cytotoxicity caused by Nanazoid on HCT-116 cells is depicted in this diagram. Nanazoid was examined at twelve different concentrations: 100, 50, 25, 12.5, 6.25, 3.125, 1.56, 0.78, 0.39, 0.19, 0.097, 0.048 ppm.

Figure 3: The average cytotoxicity caused by Nanazoid on BJ-1 cells is depicted in this diagram. Nanazoid was examined at twelve different concentrations: 100, 50, 25, 12.5, 6.25, 3.125, 1.56, 0.78, 0.39, 0.19, 0.097, and 0.048 ppm.
The inhibition concentration (IC50) of Nanazoxid on HCT-116 and BJ-1 are represented below (Table 2 and Table 3)

**Table 2. IC50 value on the HCT116 cell line**

<table>
<thead>
<tr>
<th>Drug</th>
<th>IC50</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nanazoxid</td>
<td>0.20μg/ml</td>
</tr>
</tbody>
</table>

**Table 3. IC50 value on the BJ-1 cell line**

<table>
<thead>
<tr>
<th>Drug</th>
<th>IC50</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nanazoxid</td>
<td>25.082μg/ml</td>
</tr>
</tbody>
</table>

Selectivity index = \( \frac{25.082}{0.20} = 125.41 \).

**B. Gene Expression analysis:**

By evaluating the amount of expression of Caspase-3, Bax, P53, and Bcl-2 genes in HCT-116 cells, the influence of IC50 (0.20 μg/ml for HCT116) Nanazoxid on gene regulation was examined using qRT-PCR. Gene expression changes were computed using the 2-ΔΔCt method and are shown as fold changes.

Equation: \( 2\Delta\Delta C_t = 2^{C_t \text{ (treated cells)}} - C_t \text{ (control cells)} \)

After comparing the expression levels of the Caspase-3, Bax, P53, and Bcl-2 genes in treated and untreated cells, it was discovered that the Bcl-2 gene to be slightly down by 0.169 folds, the Bax gene was over expressed by 10.72 folds, P53 was down expressed by 6.52 folds, and Casp3 was down expressed by 7.57 folds. Changes in the expression levels of the respective genes in HCT116 are depicted (Table 4 and Figure 4).Because it is a highly conserved gene that is often employed as an internal control for gene expression research, the β-actin gene was chosen as an internal control "Housekeeping gene." (Roh et al., 2000)

**Table 4. Effects of Nanazoxid on Caspase-3, Bax, p53 and Bcl-2 gene expressions in HCT116 cell line**

<table>
<thead>
<tr>
<th>Genes</th>
<th>Change in Expression (Folds)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Caspase-3</td>
<td>7.57</td>
</tr>
<tr>
<td>Bax</td>
<td>10.72</td>
</tr>
<tr>
<td>p53</td>
<td>6.52</td>
</tr>
<tr>
<td>Bcl-2</td>
<td>0.169</td>
</tr>
</tbody>
</table>

**Figure 4:** Caspase3, Bcl-2, P53, and Bax gene expression profiles are schematically shown. Normal and untreated cells were compared in gene expression levels under the same conditions.Bcl-2 gene expression was reduced by 0.169 folds, Bax gene expression increased by 10.72 folds, P53 by 6.52 folds, and Casp3 by 7.57 folds.
C. Apoptosis induced by Nanazoxid observed using Flowcytometry:

1. After the HCT116 cells were treated with Nanazoxid, there was a cell growth arrest at G\textsubscript{0}/G\textsubscript{1} phase and S phase. An increase in Pre G\textsubscript{1}, G\textsubscript{0}/G\textsubscript{1} and decrease in G\textsubscript{2}-M and slightly increase in S phase (Table 5 and Figure 5).

Table 5. After a 24-hour treatment with Nanazoxid, the cell cycle in HCT116 cells was examined

<table>
<thead>
<tr>
<th>Sample data</th>
<th>Results DNA content</th>
</tr>
</thead>
<tbody>
<tr>
<td>code</td>
<td>IC50 ug/ml</td>
</tr>
<tr>
<td>Treated</td>
<td>51.23</td>
</tr>
<tr>
<td>Control</td>
<td>48.04</td>
</tr>
</tbody>
</table>

Figure 5: The cell cycle profiles of the treated and control HCT116 cells are depicted schematically. In the G0-G1 and S phases, the proportion of cells grew, fell in the G2-M phase, and increased in the pre-G1 phase.

2. When a cell enters apoptosis, the apoptosis marker phosphatidylserine is exposed to the plasma membrane’s surface. To evaluate Nanazoxid’s apoptotic capacity in HCT116 cells, Annexin V, a Ca\textsuperscript{2+}-dependent phospholipid-binding protein with an affinity for PS, is used to identify this signal. According to the flowcytometry, the drug triggered 1.86% early, 27.44% late apoptosis and 18.01% necrosis (Table 6 and Figure 6).

Table 6. Apoptotic effects of Nanazoxid on the HCT116 cells

<table>
<thead>
<tr>
<th>Apoptosis</th>
<th>conc</th>
<th>Total</th>
<th>Early</th>
<th>Late</th>
<th>Necrosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treated</td>
<td>47.31</td>
<td>1.86</td>
<td>27.44</td>
<td>18.01</td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>1.69</td>
<td>0.46</td>
<td>0.14</td>
<td>1.09</td>
<td></td>
</tr>
</tbody>
</table>
DISCUSSION

Different antibiotics were tested in this research to see if they had anticancer action against a colorectal cancer cell line. One of the 10 drugs has anti-proliferative properties. The antibiotics with the most promising inhibitory effect were chosen for additional testing in order to figure out how these effects work.
On HCT-116 cancer cell line, benzobiotoic and quinabiotic had mild anticancer effects.

Nanazoxid had a high level of cytotoxicity in the HCT-116 cell line (IC50 of 0.20 μg/ml), but only had a little effect on normal BJ-1 cells. This means Nanazoxid is only cytotoxic to colorectal cancer cells.

The distribution of cells in the cell cycle was changed by Nanazoxid, according to the cell cycle analysis. The proportion of cells in the G0-G1 and S phases increased, while the number of cells in the G2-M phase reduced and the pre-G1 phase grew slightly. These findings point to a disruption in the cell cycle, with the cell trapped in the G0-G1 and S phases.

Nanazoxid produced 1.86 % early apoptosis and 27.44 % late apoptosis, according to flow cytometry data (Table 6). Furthermore, it caused necrosis in 18.01 % of the treated cells against 1.09 % of untreated cells. Nanazoxid had a great potential to induce apoptosis. Induction of necrosis is also somewhat elevated. As a result, the drug is being examined as a potential anticancer medication.

The treatment of HCT-116 cells with Nanazoxid resulted in a shift in the anti-apoptotic/pro-apoptotic balance of the cells toward undergoing apoptosis, demonstrating that the drug can shift the cells’ anti-apoptotic/pro-apoptotic balance toward undergoing apoptosis (Table 6).

The analysis indicates that there was a very little down expression in the Bcl-2 gene by 0.169 folds in the treated and untreated cells, overexpression in the Bax gene by 10.72 folds, P53 by 6.52 folds, and Casp3 by 7.57 folds in the treated cells. (Table 4). Nanazoxid was shown to have a strong potential to trigger apoptosis by downregulating Bcl2 genes and boosting the overexpression of pro-apoptotic proteins like Bax and apoptotic proteins like P53 and Casp3, making it a prospective anticancer drug.

CONCLUSION

Nanazoxid has a strong influence on cell proliferation and apoptosis, and it has been shown that it affects the cell cycle and gene expression of Casp3, Bax, P53, and Bcl-2. However, more research into the molecular mechanism that controls the balance between cellular growth and death is needed. Nanazoxid’s ability to generate high cytotoxicity in a colorectal cancer cell line in vitro. Because of its great anticancer activity, it should be explored further to see if it may be utilized as a treatment for colorectal cancer. Benzobiotoic and Quinabiotic can also be studied further because they have a mild anticancer impact.

REFERENCES


المخلص العربي

نشاط عقار نانازوکسید كمضاد للسرطان في المختبر ضد خلايا سرطان القولون والمستقبل ومساراته الجزئية

نورهن محمد عاد ١، سلوى محمد الهلوطي ٢، عماد محمد الزيات ٣، أحمد عيد الشريف ٤

١ - طالب ماجستير في قسم البيوتكنولوجيا، كلية العلوم، جامعة القاهرة، القاهرة، مصر.
٢ - معمل زراعة الخلايا الحيوية، قسم الخلية، المركز القومي للبحوث، الدقي، الجيزة، ١٢٣٢، مصر.
٣ - قسم علم الحيوان، كلية العلوم، جامعة القاهرة، القاهرة، مصر.
٤ - قسم الكيمياء، كلية العلوم، جامعة القاهرة، القاهرة، مصر.

تكتب إعداداً اتّفق الإدراة قصرية كفخ إلك للعلاج. فالنهاية قريبة كناب لازم للاستقدام بلزن كمزة من أهداف جزئية جديدة لعقار مرخص سريّاً معروفاً في مجموعة متنوعة من الأمراض، سواء الموجودة أو الجديدة. أظهرت بعض الدراسات المبكرة أن هيروكسي كلوروكين، الذي يستخدم من ذه منظم ميل جزئية لعلاج الملاريا، يستخدم لتقليل الحمى والتّت ارضاً كودف ١٩٩٥.

السرطان هو مشكلة صحية رئيسية في جميع أنحاء العالم. خلال السنوات الماضية، ثبت أن بعض الأمور الأخرى هو دورها مضادة للسرطان مثل الكشفات الفراغ الحيوية في الذين أسربين تذكير سرطان القولون والمستقبل وفقاً للفحوصات. وفقاً للمعهد الوطني للسرطان فإنثاً كرآة سرطان القولون والمستقبل سيقوم باكتشاف أكثر أنواع السرطانات انتشرت في مصر، حيث يُقدر٪ ٤٤٪ من السرطانات. كان من المتوقع أن يزيد عدد ضحايا سرطان القولون (باستثناء سرطان القولون المستقيم) إلى حد ما عن ١٠٠٠،٠٠٠ هنا في هذا المشروع وهو إعادة استخدام مضادات الحيوية المستخدمة سريّاً في السوق المصري لسأح القولون والمستقبل. تم اختبار فعالية مضادة للسرطان للعديد من مضادات الحيوية من خلال فحصها في ٢، أظهرت النتائج أن دواء واحد كنهاية مضاد للسرطان خاصة سرطان القولون والمستقبل مع Nanazoxid كدواء واحد كنهاية مضاد للسرطان خاصة سرطان القولون والمستقبل مع IC50 (٢٠٠ ميكروغرام / مل) وتم اختياره لموارد اقتصاد على الخلايا البشرية خط ورغب للتحليل الجزئي. نتيجة لذلك، أثبت دواء مضاد حيوي وهو Nanazoxid كدواء مضاد حيوي وهو آمن يمكن إعداده كعلاج مضاد للقولون والمستقبل مع مزيد من البحوث والتطوير الم Mansion.

الكلمات المفتاحية: سرطان القولون والمستقبل - إعادة استخدام الإدراة - المضادات الحيوية - خط الخلايا البشرية - الأسبرين - كودف ١٩٩٥ - نانازوکسید.