Intensification of resveratrol cytotoxicity, pro-apoptosis, oxidant potentials in human colorectal carcinoma HCT-116 cells using zein nanoparticles

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Resveratrol (RSV), a non-flavonoid stilbene polyphenol, possesses anti-carcinogenic activities against all the major stages of cancer. Zein nanoparticles (ZN NPs) have been utilized successfully in delivery of variant therapeuticals by virtue of their histocompatible nature. The goal of this work was to comparatively explore the anti-proliferative, pro-apoptotic and oxidative stress potentials of RSV-ZN NPs versus RSV against human colorectal carcinoma HCT-116 cells. ZN-RSV NPs were developed and assayed for particle size analysis and RSV diffusion. The selected formula obtained 137.6 ± 8.3 nm as mean particle size, 29.4 ± 1.8 mV zeta potential, 92.3 ± 3.6% encapsulation efficiency. IC50 of the selected formula was significantly lower against HCT-116 cells versus Caco-2 cells. Also, significantly enhanced cellular uptake was generated from RSV-ZN NPs versus free RSV. Enhanced apoptosis was concluded due to increased percentage cells in G2-M and pre-G1 phases. The pro-apoptotic potential was explained by caspase-3 and cleaved caspase-3 increased mRNA expression in addition to NF-κB and miRNA125b decreased expression. Biochemically, ZN-RSV NPs induced oxidative stress as demonstrated by enhanced reactive oxygen species (ROS) generation and endothelial nitric oxide synthase (eNOS) isoenzyme increased levels. Conclusively, ZN-RSV NPs obtained cell cycle inhibition supported with augmented cytotoxicity, uptake and oxidative stress markers levels in HCT-116 tumor cells in comparison with free RSV. These results indicated intensified chemopreventive profile of RSV due to effective delivery utilizing ZN nano-dispersion against colorectal carcinoma HCT-116 cells.

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Resveratrol (trans-3',4',5'-trihydroxystilbene; RSV) is a natural occurring stilbene polyphenolic chemopreventive and anticancer drug. RSV can be naturally obtained daily via grapes, blueberries, raspberries, mulberries, and peanuts. RSV has been sorted as a natural lipophilic phytoalexin for being synthesized by plants due to infection, fungal attack, ozone exposure and UV exposure. RSV was first isolated by the Japanese scientist Mitchio Takaoka from Veratrum grandiflorum O. Loes. in 1939. RSV belongs to a vast group of physiologically active compounds that exhibit a wide variety of therapeutic effects including anti-oxidant, anti-inflammatory, anti-atherogenic, anti-fungal, cardioprotective, anti-atherosclerosis and anti-tumor effects. RSV was first introduced in the early nineties in spite of over 20,000 research papers and around 130 human clinical trials interested about RSV. RSV is also reported to suppress oxidative stress affecting nitric oxide generation that affects DNA damage, cell cycle, apoptosis, and proliferation. Specifically, RSV enhanced apoptosis of HCT-116 cells via variant molecular mechanisms.

**Colorectal carcinoma (CRC) is a universal leading cause of death in men (10%) and women (9.2%) and ranks the third most common cancers and the fourth most common cause of cancer-related death.** Globally, CRC mortality will be increased by 56% between 2020 and 2040. In 2040, the estimated number of deaths due to the disease is 1.6 million deaths worldwide according to the international Agency for Research on Cancer (IARC). The elevated percent of CRC patients can be related to increased ageing population, non-healthy dietary habits and an increase in risks of smoking, obesity and low physical exercise.

The obtained resistance to tumors due to conventional therapeutics is a crucial incident in cancer treatment. Particularly, apoptosis resistance enables tumors cells evade innate surveillance mechanisms and disturbing intracellular signaling cascade leading to tumor promotion and metastasis. Distraction of apoptosis pathways may shift cancerous cells to be treatment resistant beside tumorigenesis promotion. Induction of apoptosis by anti-carcinogenic molecules has been conformed with tumor response. Treatment resistance was reported to be as a result of apoptosis potential failure especially when apoptosis is the main action of anticancer mechanism. Apoptosis enhancement is a site of research interest to support cancer clinical management. Induction of apoptosis in tumor cells is a frontier strategy with limit concurrent damage to normal cells.

Nanoparticulate delivery systems are characterized by eminent capability of enhancing drug solubilization, sustained circulation time, targeting at tumor via augmented permeability and retention effects, enhanced overcoming of intratumoral heterogeneity in cancerous cells due to surface receptor expression and reduction of antitumor molecule toxicities and dose as a result of improving cytotoxicity and cellular uptake. Zein nanoparticles (ZN NPs) are highly suggested nanocomposites because of their plant origin and appropriate method of fabrication. ZN is obtained from maize plant. According to United States Food and Drug Administration, ZN is Generally Recognized as Safe (GRAS), biocompatible and inexpensive excipient that used ordinarily in food industry, pharmaceutical industry, vaccine delivery, tissues engineering and biomedical applications. Also, ZN has elevated thermal resistance and good oxygen barrier capability for the loading of thermosensitive compounds. Pharmaceutical nanotechnology enhances antitumor therapeutics profiles. The aim of this study was to comparatively investigate the antiproliferative, pro-apoptotic and oxidative stress profiles of RSV-ZN NPs in comparison to free RSV in human colorectal carcinoma HCT-116 cells.

**Results and discussion**

RSV was studied extensively along last decade for obtaining cancer therapeutics promising outcomes in both cancer prevention and treatment. The anticancer molecular mechanism is still not elucidated. Effective formulation of RSV was frequently faced the difficulty of very low oral bioavailability due to reduced water solubility and low photostability. Just less than 1% bioavailability was reported due to severe metabolism in liver and intestines. Pharmaceutical nanotechnological techniques offer successful approach to overcome this obstacle. Nanostructures of RSV are highly recommended to increase water solubility, facilitate biological membranes cross, ameliorate absorption and stability and enhance pharmacokinetic parameters and bioavailability. This will have significant and promising effect on therapeutic potential of RSV to ameliorate its cytotoxicity and pro-apoptotic activity against HCT-116 cells. The efficiency of nanostructures loaded with RSV will extended to overcome the suggested multiple drug resistance in various tumors and minimize toxicity via enhancing permeation and retention effect of nanostructures.
Characterization of ZN-RSV NPs. RSV Nanocomposites were fabricated in three variant formulae; R1, R2 and R3. Table 1 displayed formulation characteristics for all developed formulae. The most reduced mean particle size was related to formula R1 which was 137.6 ± 8.3 nm. This obtained mean particle size of ZN-RSV NPs is suggested to enhance absorption and residence time inside biological systems. As shown, particle size reduction was directly related to elevation of ZN content through the prepared formulae. The higher particle size of R3 formula could be generated due increased quantity of RSV on nanosphere surface, which could lead to maximized interfacial tension between particle surface and aqueous medium, so consequently particle diameter will be enlarged. Also, R1 formula obtained the highest zeta potential magnitude (29.4 ± 1.8 mV) and the least polydispersity index magnitude (0.19 ± 0.01). Elevated zeta potential value reflects nanostructures dispersion stability as it will withstand aggregations because of charge of ZN surface. R1 formula obtained 92.3% ± 3.6 encapsulation efficiency. The enhanced EE by increasing of in ZN amount in formula could be related to RSV partitioning to the hydrophobic matrix of ZN. The elevated encapsulation efficiency by elevation in ZN amount in formula might be due to integrated partitioning of RSV to ZN hydrophobic matrix which reflects a relation between RSV binding affinity and ZN polymerization degree. The ability of nano-delivery systems to incorporate high percent of payloads was previously reported. The method utilized for fabrication was adequate in obtaining high encapsulation efficiency values. This could be explained regarding to the natural unfolding of ZN molecules at alkaline pH which offers more reactive sites for maximized crosslinking and also remarkable reduction of void spaces inside ZN nanospheres which will positively contribute to ameliorated bioavailability, delivery and consequently the therapeutical response.

ZN-RSV NPs formula diffusion is presented in Fig. 1. R1, R2 and R3 formulae displayed sustained RSV diffused profile, which obtained magnitudes of 94.1 ± 3.6%, 90.3 ± 4.5 and 84.4% ± 6.9 after 36 h, respectively. R1 formula was chosen for further investigations. Sustaining RSV diffusion could be attributed to ZN nature hydrophobic structure. ZN generates a delay in the water penetration which could be attributed to the measured RSV sustained diffusion. RSV diffusion exhibited biphasic sustained permeation profile. The diffusion displayed initial burst effect related to rapid release of trapped RSV to direction of ZN NPs surfaces. After initial burst, a reduction in diffusion rate was attained as RSV has to penetrate longer path in core matrix to completely release from nanospheres. Also, RSV diffusion rate was affected by water intake rate. The prolonged release of RSV due to ZN NPs will support the pharmacokinetic potential due to short half-life and reported irregular profile in human after oral dose of RSV by that obtained two peak plasma concentration after 1 h and 6 h due to enteric recirculation.

The particles morphology of R1 formula was examined by SEM imaging and showed compactly dispersed spheres with smooth surfaces. The diameter of scanned spheres was consistent with particle size measured by laser diffraction technique. The noticeable compactness of nano-particulates may be related to freeze drying process.

Serum stability. ZN-RSV NPs colloidal stability was estimated by monitoring the changes in sample mean particle size (Fig. 3). After incubation in FBS, ZN-RSV NPs displayed initial elevation in its mean particle size in the first 15 min and then decreased rapidly to starting value with non-significant difference. The developed RSV nanocomposites attained satisfied pattern in FBS which suggest similar profile in in vivo studies due absence.
of persistent aggregation due to possible interacting with variant molecules inside the biological environment around nanocomposites.

**Cytotoxicity.** Cytotoxicity of the different treatments using mitochondrial function (MTT reduction) results are exhibited in Fig. 4. It indicates distinguished augmentation of ZN-RSV NPs antiproliferative activity against HCT-116 cells. ZN-RSV NPs recorded enhanced activity of about \( IC_{50} = 4.15 \, \mu g/ml \) compared to free RSV (46 \( \mu g/ml \)). This premium cytotoxicity revealed the potential of ZN NPs due to noticeable reduced particle size of prepared nano-particulates that can interact with biomolecules within cells or on their surfaces, in addition to adequate encapsulation inside ZN NPs and complete diffusion of RSV that enhance potentiate cytotoxicity\[^{52,53}\].

Also, it was very noticeable the cytotoxic effect of ZN alone in HCT-116 cell culture. ZN cytotoxicity may explain the obtained enhanced anti-proliferative action of ZN-RSV NPs. Cytotoxicity was enhanced in similar studies utilizing ZN nanoparticles as reported for dual delivery of exemestane and RSV\[^{54}\], Piceatannol\[^{55}\] and pterostilbene\[^{56}\]. Similar studies reported highlighting cytotoxicity of ZN against several types of cancers cells\[^{44}\].
Conclusively, ZN represented an optimistic biomaterial for fabrication of RSV nano-formulations and also cytotoxicity ameliorating.

**Cellular uptake.** Figure 5 exhibited that free RSV cellular uptake obtained 19.1 ± 1.3% and 31.4 ± 2.7% at 2 and 4 h of incubation, respectively. Significantly elevated cellular uptakes were observed with ZN-RSV NPs incubations, which reached 41.6 ± 23.1% and 62.3 ± 5.2% at after 2 and 4 h of incubation, respectively. The results indicated ameliorated cellular uptake significantly by ZN-RSV NPs compared to free RSV. This also confirmed the ability of ZN NPs to enhance HCT-116 cells uptake of RSV that may ameliorate pro-apoptotic profile. The obtained results concluded significant ameliorated RSV cellular uptake due to the developed formula. This confirmed the ability of ZN NPs to enhance HCT-116 cells uptake of RSV which also confirm the efficacy of ZN improved Enhanced Permeability and Retention (EPR) cellular uptake in tumor microenvironment. The cellular uptake of polymeric nanoparticles which possess < 200 nm mean diameter is described by an endocytotic action. Also, positively charged nanoparticles generated a higher rate of membrane internalization in comparison with negatively charged ones due to enhanced adsorptive-mediated transcytosis catalyzed by electrostatic interaction yielded between positively charged structures of ZN and negatively charged cells membranes.

**Cell cycle progression.** Rapid growth properties were exhibited from control untreated HCT-116 cells. According to Fig. 6a, it obtained 52.06 ± 3.7% at the G0/G1 phase, 39.08 ± 2.9% at the S phase, 8.06 ± 0.4% at the G2-M phase, and 1.78 ± 0.6% at the pre-G1 phase. Other RSV, ZN and ZN-RSV NPs incubations displayed down the proliferation of HCT-116 cells, especially in the G0/G1 and S phases (Fig. 6b–d). Regarding to accumulation of cells in the pre-G phase, it records 1.78 ± 0.6%, 23.14 ± 1.1%, 7.98 ± 0.5 and 25.72 ± 1.4% of control HCT-116 cells, RSV, ZN, and ZN-RSV NPs groups, respectively. RSV antiproliferative potential is correlated with apoptosis induction capability and G1 phase cell cycle arrest in colon cancer cells. Also, the concluded pro-apoptotic activity was in harmony with the obtained RSV enhanced induced apoptosis in HCT-116 cells three-dimensional culture leading to formation of a luminal cavity by suppressing PDE4 activity. Figure 6e exhibited the obtained changes in cell cycle phases graphically.

**Annexin-V staining.** In order to confirm the generated cell apoptotic death, the percentage of HCT-116 cells with positive annexin-V staining were estimated in the control, RSV, ZN and ZN-RSV NPs incubations (Fig. 7a–d). ZN-RSV NPs distinctly increased the early, late, and total cell death in comparison with other incubations. A graphical explanation for cell death different types was explained in Fig. 7e. ZN-RSV NPs obtained the most functional activity in enhancing pre-G phase, which confirms apoptotic cell death. This indicated early and late apoptotic death, as well as total cell death. The obtained results confirmed the potential of RSV to induce apoptosis in colon cancer cells. Similar RSV enhanced pro-apoptotic was obtained in HT29 colon cancer cell. Also, these results support the utilization of ZN Nano-dispersed delivery systems for enhancing apoptotic and cellular uptake potential of piceatannol and pterostilbene.

**mRNA expression of apoptosis-related genes.** Figure 8 displayed the results of mRNA expression of pro-apoptotic genes (CASP3 and cleaved caspase-3) and anti-apoptotic genes (miRNA125b and NF-xb) related to ZN-RSV NPs treated HCT-116. The cells exhibited statistically variant ratios. It was concluded that HCT-116 cells treated with ZN-RSV NPs obtained the highest values of expression related to caspase-3, cleaved caspase-3, miRNA125b and NF-xb genes. Activation of caspase-3 is so fundamental for apoptosis inducement and cancer targeting obtained by anti-tumor therapeutics; thus, it is considered a main indicator for sensitivity estimation of cancerous cells to therapeutics. All treated groups (RSV, ZN and ZN-RSV NPs) exhibited folds change of 5.75, 3.54 and 7.21,
respectively. Regarding to cleaved caspase-3 expression, all treated groups (RSV, ZN and ZN-RSV NPs) exhibited folds change of 1.68, 1.43 and 2.41, respectively. ZN-RSV NPs displayed elevated apoptotic effect in comparison with untreated positive control. ZN-RSV NPs obtained 1.25-fold elevation in caspase-3 level and 1.43-fold elevation in cleaved caspase-3 level in comparison with free RSV. Caspase-3 is considerably stimulated by the death protease, activating the specific cleavage of numerous major cell proteins, and it contributes mainly in programmed cell death. A relation between RSV mechanism of action and caspase activation via blocking antiapoptotic proteins belonging to heat shock proteins was investigated.

Figure 6. Impact of ZN-RSV NPs on cell cycle phases. (a) Control, (b) RSV, (c) ZN, (d) ZN-RSV NPs, (e) graphical presentation of each phase. *Significantly different in comparison with control group at p < 0.05.
Regarding to miRNA125b expression, all treated groups (RSV, ZN and ZN-RSV NPs) displayed folds change of 0.503, 0.52 and 0.387, respectively. ZN-RSV NPs exhibited higher anti-apoptotic effect compared with untreated positive control. ZN-RSV NPs obtained 1.3-fold decrease in miRNA125b level compared with free

Figure 7. Impact of ZN-RSV NPs on annexin-V FITC positive staining HCT-116 cells. (a) Control, (b) RSV, (c) ZN, (d) ZN-RSV NPs, (e) graphical presentation of early and late apoptotic, necrotic and total cell death. *Significantly different in comparison with corresponding control group at p < 0.05.
These results confirmed the apoptotic role of RSV in cancer progression through modulation of microRNAs (miRNAs) which has not been studied. Also, colon cancer progression was related to miRNAs that activating either as oncogenes or tumor-suppressor genes. In comparison to normal tissues, miRNAs have been dysregulated in colon cancer tissues.

NF-κB pathway is the fundamental contributor to colon tumors where RSV inverts NF-κB activation, which is responsible for inducing inflammatory cytokines. The results of NF-κB expression showed folds change of 3.51, 3.8 and 4.96, respectively. ZN-RSV NPs displayed elevated anti-apoptotic effect in comparison with untreated positive control. ZN-RSV NPs attained 1.41-fold decreasing in NF-κB level in comparison with free RSV. As investigated by NF-κB, as nuclear transcription factor involved in carcinogenesis, is implicated in cell proliferation, differentiation, apoptosis, tumorigenesis and cellular stress reactions. NF-κB activation might regulate several genes expression leading to cell survival. These results were in harmony with where RSV was concluded to suppress NF-κB by blocking TNF-induced activation of NF-κB in a dose- and time-dependent manner.

Reactive oxygen species assay (ROS). As displayed in Fig. 9, ZN-RSV NPs obtained the largest green fluorescence intensity. ZN-RSV NPs obtained highest ROS activity (156.1 ± 9.4 Pg/106 million cells) which equal 1.27 upfolds of free RSV sample. This concluded that ZN-RSV NPs could induce oxidative stress in cancer cells by increasing ROS formation via oxidation of NADPH. RSV demonstrated a protective potential regarding to ROS and nitric oxide synthesis. ROS can generate subsequent toxic radicals like hydroxyl, hydrogen peroxide and superoxide which react to DNA and proteins on cellular level leading to DNA damage and lipid peroxidation. The anti-peroxidative action of RSV was reported and correlated to capability of phenolic groups to regain lipid hydroperoxyl, hydroxyl and superoxide anions. Also, apoptosis induced by RSV in NCI-H460 non-small cell lung cancer cells and human colon cancer cells was contributed to ROS levels regulations in that trigger downstream signaling pathways as antioxidant or pro-oxidant.

eNOS activity assay. As reported eNOS is expressed in many cell lines as colon cell line HCT-116. The response was explained as a response of tumor cells to the therapeutic agent. The results revealed increased levels of eNOS isoenzyme in ZN-RSV NPs sample in comparison with control and free RSV samples as exhibited in Fig. 10. There was increased significant production of eNOS in RSV and ZN-RSV NPs sample with obtained values of 10.72 and 12.58 ng/million cells, respectively. Zein base nano-particulates was a successful option for enhancing NO production induced by RSV in HCT-116 cells. This was in accordance with where suppression of tumorigenicity and abrogation of cancer metastasis is related directly to direct NO production, which confirm apoptosis potential. Also, it has been concluded that RSV enhanced endothelial NO release through long and short-term effects where rapid effect comprises phosphorylation of eNOS by AMP-activated kinase, or extracellular signal regulated kinase 1 and 2, and deacetylation of eNOS by sirtuin.
**Materials and methods**

**Materials.** Zein (CAS number: 9010-66-6, ZN), RSV (CAS number: 501-36-0) and Absolute Ethanol were purchased from Sigma-Aldrich (St. Louis, MO, USA). All other solvents and chemicals used were of analytical grade.

**ZN-RSV NPs formulation.** ZN-RSV NPs were prepared using nanoprecipitation method[^40][^42][^55]. Three ZN-RSV NPs formulae (R1–R3) were prepared and varied according to the applied drug:polymer ratios. RSV and ZN were dissolved in 10 ml of 85% ethanol using of vortex (Velp scientifica, ZX3, Usmate, Italy) and ultrasonic probe (Vibra-Cell VCX750; Sonics and Materials Inc., Newtown, CT, USA). The ethanolic dispersion was flowed into deionized water and magnetically stirred at room temperature for 3 h at 2000 rpm to vaporize content of ethanol. The aqueous suspension was then centrifuged at 20,000×g speed then lyophilized using cryoprotectant of trehalose.

**ZN-RSV NPs characterization.**

**Particle size analysis.** Laser diffraction technique was utilized for particle size analysis of ZN-RSV NPs formulae (R1–R3). Using disposable cuvette, one milliliter of the sample was diluted in deionized water. The average particle size was then determined.

**Encapsulation efficiency (EE).** Ethanol solubilized samples were filtered through 0.22 μm filters, and analyzed for (EE) using reported high-performance liquid chromatography (HPLC) (Agilent 1200, Agilent Technologies, Santa Clara, CA, USA) equipped with a C18 column (5 μm, 4.6 mm×250 mm), photodiode array detector (PDAD) at 306 nm (Waters, Milford, USA), adjusted column temperature at 35 °C, mixture of methanol and purified water (50:50 v/v) as mobile phase and 1 ml/min flow rate according to[^82]. RSV EE was estimated according to the following equation:

\[
EE\,\% = \frac{\text{amount of RSV in the nanocomposites}}{\text{amount of RSV initially added}} \times 100. \tag{1}
\]

![Figure 9](https://www.nature.com/scientificreports/)  **Figure 9.** ROS activity in control, RSV and ZN-RSV NPs samples. *Significantly different in comparison with control group at p < 0.05. †Significantly different in comparison with RSV group at p < 0.05. ‡Significantly different in comparison with ZN group at p < 0.05.

![Figure 10](https://www.nature.com/scientificreports/)  **Figure 10.** Levels of eNOS in control, RSV and ZN-RSV NPs samples. *Significantly different in comparison with control (p < 0.05). †Significantly different in comparison with RSV (p < 0.05).
**In vitro RSV diffusion.** RSV diffusion was carried out according to\(^\text{42}\) using automated Franz diffusion cell apparatus (MicroettePlus, Hanson Research, Chatsworth, CA, USA) of 7 ml chamber volume and 1.76 cm\(^2\) diffusion area using Phosphate-buffered saline (0.01 M, pH 7.0) at maintained temperature of 37 ± 0.5 °C. Synthetic nylon membrane of 0.45 μm pore size (Pall Corporation, Port Washington, NY, USA) was the utilized diffusion membrane. The study was proceeded for 36 h with selected intervals at 0.25, 0.5, 1, 2, 4, 8, 12, 18, 24 and 36 h. Aliquots were analyzed for RSV content utilizing validated HPLC method\(^\text{82}\).

**Nanoparticles morphology.** Scanning Electron Microscope (SEM) instrument (JEM 100-CX; JEOL, Tokyo, Japan) was used to investigate nanoparticles surface morphology of ZN-RSV NPs. Using the lyophilized powder, sample was coated under vacuum by gold after fixation.

**Serum stability.** ZN-RSV NPs Colloidal stability was examined in Fetal Bovine Serum (FBS, Gibco, Thermo Fisher Scientific, MA, USA)\(^\text{56}\). 1 ml of 70% FBS was added to 200 µl of sample suspension and then incubated at 37 °C for 48 h followed by magnetic stirring at 600 rpm. At selected time intervals of incubation, particle size analysis was carried out.

**Cell lines and culture.** Human colorectal carcinoma cell line (HCT-116), Human colorectal adenocarcinoma (Caco-2), Human colorectal adenocarcinoma (HT-29) and Human somatic cell hybrid (EA.hy926) cells were supplied from Nawah Scientific, Cairo, Egypt that were purchased from the American Type Culture Collection (ATCC; Manassas, VA, USA). HCT-116 were cultured in ATCC-formulated McCoy’s 5a Medium Modified and Caco-2, HT-29 and EA.hy926 Cells were cultured in ATCC-formulated Eagle’s Minimum Essential Medium. The mediums were supplemented with fetal bovine serum to a final concentration of 10% and 20%. Culture plates and flasks (SPL Life Sciences, Korea) were subjected to addition of Penicillin (100 U/ml) and streptomycin (100 mg/ml) previously. Cells were maintained inside carbon dioxide incubator at 37 °C in a humidified atmosphere containing 5% CO\(_2\) (Thermo Electron Corporation, Forma series II, 3141, USA) to keep the cells in a sub-confluent state.

**Cytotoxicity assay.** MTT assay was utilized to evaluate the antiproliferative activity using the kit of ABCAM, Cambridge, UK according to\(^\text{51}\), where HCT-116, Caco-2, HT-29 and EA.hy926 cells were introduced into 96-well plates (TPP, Switzerland) as 2 × 10\(^3\) cells/well approximately. Wells were incubated with RSV, equivalent weight of ZN or ZN-RSV NPs using a range of concentrations at logarithmic manner at 37 °C in a carbon dioxide incubator for 48 h.

**Cellular uptake analysis.** In the presence of 5% CO\(_2\), HCT-116 cells were incubated overnight (1 × 10\(^5\) cells/dish) at 37 °C for 2 and 4 h after treating with ZN-RSV NPs IC\(_{50}\) value, equivalent concentration of RSV or ZN. Lysis solution was added for 30 min at 37 °C after monolayers were washing. HPLC technique was utilized for analyzing of cell lysates aliquots.

**Cell cycle progression analysis.** This assay was carried out as described by\(^\text{55,56}\) using the same reagents and kits.

**Annexin-V assay.** This assay was carried out as described by\(^\text{55,56}\) using the same reagents and kits.

**mRNA expression of apoptosis-related genes by quantitative real-time polymerase chain reaction (RT-PCR).** This assay was carried out as reported by\(^\text{55,56}\) using the same reagents and kits. Table 2 gives the primer sequences for caspase-3, miRNA125b, NF-xB and β-actin. Results were validated using the relative quantification (ΔΔCT) method. The genes expression was estimated in triplicates and runs mean was normalized with β-actin mean.

**Reactive oxygen species assay (ROS).** IC\(_{50}\) concentrations of ZN-RSV NPs and equivalent concentrations of RSV and ZN were incubated in 96 well plates with adjusted cell density at 5 × 10\(^3\) HCT-116 cells/well for period

| Primer     | Sequence                                                |
|------------|---------------------------------------------------------|
| Caspase-3  | Forward primer 5′-GTGGAACGTGACGATATGGC-3′               |
|            | Reverse primer 5′-CAGAAATGTACCCTGAAACC-3′               |
| miRNA125b  | Forward primer 5′-CTTGCCAGAAACGCTACATGG-3′              |
|            | Reverse primer 5′-GTGCAATCGACATACCTG-3′                 |
| NF-xB      | Forward primer 5′-ATCCCATCTGGACAAAACGG-3′               |
|            | Reverse primer 5′-CTGTGCCTGGATAATAACCT-3′               |
| β-Actin    | Forward primer 5′-AAGATCCTGGACGAGGCTGG-3′               |
|            | Reverse primer 5′-CAGCAGCTGTGTTGACAGAGG-3′              |

Table 2. Primer sequences used for RT-PCR.
of 24 h. For staining, we used 10 μM 2,7-dichlorofluorescein diacetate (DCFDA) for 45 min. The used kit was ABCAM, Cambridge, UK.

**Endothelial nitric oxide synthase (eNOS) activity assay.** Enzyme-linked immunosorbent assay kit (RayBio’ Human eNOS ELISA Kit, Norcross, GA, USA) was used to measure eNOS levels using 96-well plate. After adding of standards and samples into plates, biotinylated antihuman eNOS antibody is added where eNOS was present bound to plate wells by immobilized antibody. HRP conjugated streptavidin is introduced to wells after washing. Second washing is carried out before the addition of TMB substrate solution. Then color was generated according to quantity of eNOS attached. Color was changed from blue to yellow after addition of stop solution. Finally, color intensity was measured at 450 nm.

**Statistical analysis.** Study data were displayed as means ± standard deviation (SD) of at least three independent runs. Comparative statistics between treatments was carried out by Student’s t test or one-way analysis of variance followed by Tukey’s test (criterion of significance was considered two-tailed p value less than 0.05). The utilized software was IBM SPSS software (Trial online version, SPSS Inc., Chicago, IL, USA).

**Conclusions**

The present study explored the ameliorating impact of nanoformulation on the molecular mechanism of RSV as anticancer molecule against Human CRC cells HCT-116. ZN-RSV NPs were fabricated with mean particle size of 137.6 ± 8.3 nm using the promising biomaterial of ZN. The developed ZN-RSV-NPs attained elevated EE%, 36 h sustained diffusion and colloidal stability. The enhanced cytotoxicity and cellular uptake, ameliorated pro-apoptotic potential supported by enhanced mRNA expression of caspase3, cleaved caspase-3, miRNA125b and NF-kb genes confirmed the enhanced anti-proliferative and pro-apoptotic potentials of ZN-RSV NPs. Biochemical examining exhibited increased generation of ROS and eNOS levels due to the ZN-RSV NPs formulation which displayed enhanced oxidant profile. The obtained results concluded therapeutic intensification of RSV antitumor potential against Human CRC cells HCT-116 due to the developed nano-delivery system.

**Data availability**

The datasets generated and analyzed during the current study are available from the corresponding author on reasonable request.

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A.M.H., M.M.S., M.T.K., A.B.A. and D.F.T. designed and conducted the experiments. R.F.T., M.M.S., M.A.Z., H.I.K., A.B.A. and D.F.T. analyzed the results. All authors wrote and reviewed the manuscript.

Competing interests
The authors declare no competing interests.

Additional information

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