Cytotoxic Property of *Grias neuberthii* Extract on Human Colon Cancer Cells: A Crucial Role of Autophagy

Luis M. Guamán-Ortiz,1 Juan C. Romero-Benavides,2 Alirica I. Suárez,2,3 Stephanie Torres-Aguilar,1 Paola Castillo-Veintimilla,2,4 Jimmy Samaniego-Romero,1 Kevin Ortiz-Díaz,1 and Natalia Bailon-Moscoso

1Departamento de Ciencias de la Salud, Universidad Técnica Particular de Loja, Loja 1101608, Ecuador
2Departamento de Química y Ciencias Exactas, Universidad Técnica Particular de Loja, Loja 1101608, Ecuador
3Facultad de Farmacia, Universidad Central de Venezuela, Caracas 1050, Venezuela
4Programa Nacional para el Abordaje Multidisciplinario de las Parasitosis Desatendidas en el Ecuador PROPAD, Instituto Nacional de Investigaciones en Salud Pública LIP, Guayaquil 3961, Ecuador

Correspondence should be addressed to Natalia Bailon-Moscoso; ncbailon@utpl.edu.ec

Received 28 June 2019; Accepted 10 March 2020; Published 1 April 2020

1.Introduction

Colorectal cancer ranks third in terms of incidence, but second in terms of mortality. [1] Colorectal cancer is increasing in Central and South America due to an ongoing transition towards higher levels of human development. [2] Understanding the mechanisms underlying the effect of apoptotic/autophagic regulators should generate new ideas and opportunities for chemotherapeutic intervention and the potential treatment of cancer patients. [3].

Natural products continue to be an important source of leads for new medicines. Historically, natural products from plants and animals have been the source of virtually all medicinal preparations and, more recently, natural products have continued to enter clinical trials or to provide leads for compounds that have entered clinical trials. [4] The production of secondary metabolites is favored by the different microenvironments. Continental Ecuador is a region with the third-highest density of endemic plant species worldwide. It is shown in [5] that *G. neuberthii* (Lecythidaceae) is endemic to Colombia, Ecuador, and Peru. In accordance with the ethnomedical uses reported in various herbaria from Ecuador and bibliographical references, medicinal uses (including antitumor) described for *G. neuberthii* are related

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This is a research article discussing the cytotoxic property of the *Grias neuberthii* extract on human colon cancer cells, focusing on the role of autophagy. The study includes experiments on two colon carcinoma cell lines, RKO (normal p53) and SW613-B3 (mutated p53), and identifies three chemical compounds responsible for the observed effects.
2. Materials and Methods

2.1. Plant Material. *G. neuberthii* was collected on a farm in Lumbaqui (00°01′46″ Lat. S; 77°10′24″ Long. O, 366 m.a.s.l.) Sucumbios Province of Ecuador. A sample specimen (LOJA-49) was deposited in the Herbarium of Universidad Nacional de Loja, Ecuador, and identified by Xavier Cornejo and Zhofre Aguirre.

2.2. Preparation Extract. The aerial parts (leaves, stem bark, fruit, and seed) were reduced to fine particles by grinding to a suitable size and then were dried at 30°C for seven days in dryer trays with air flow.

The dried and ground aerial parts of *G. neuberthii* (4045 g) were macerated at room temperature for 72 h in a light-free environment, with hexane, ethyl acetate, and methanol, sequentially, with 5 L of each solvent; the procedure was repeated three times. The extracts were filtered with an isocratic system of hexane, ethyl acetate, and methanol, sequentially, with 5 L of each solvent; the procedure was repeated three times. The extracts were filtered with a rotary evaporator (Buchi R210, Switzerland), and subsequently stored at 4°C and protected from light until further use.

Thin-layer chromatography using aluminum plates coated with silica gel 60 F254 (Merck, Germany) was performed on each extract.

For biological studies, stock solutions (40 mg/mL) were prepared in dimethylsulfoxide (DMSO—Sigma Aldrich, USA) and stored at -20°C until use. The aliquots were diluted to obtain the appropriate concentrations before use.

2.3. Phytochemical Screening. Phytochemical screening to test for the presence of secondary metabolites (alkaloids, terpenoids, steroids, flavonoids, tannins, saponins, and quinones) and proteins, carbohydrates, and fats and oils in the extracts was carried out using standard procedures [8].

2.4. Characterization and Identification of Secondary Metabolites. Melting points were determined using a Fisher-Johns apparatus. The 1H and 13C NMR spectra were recorded at 400 MHz and 100 MHz, respectively, on Varian 400 MHz Premium Shielded Equipment (Varian, USA) using tetramethylsilane as an internal reference. CDCl3, CD3OD, and DMSO-d6 were used as solvents; chemical shifts were expressed in parts per million (ppm), and coupling constants (J) were reported in Hz.

2.5. Extraction and Isolation of Compounds. The most active extract, methanol extract of stem bark (GNSBM), was partitioned, and 30 g was dissolved in MeOH::H2O (9:1) in a ratio of 1:20 (extract: solvent) (540 mL of MeOH and 60 mL of H2O) and sequentially partitioned three times with 400 mL of each solvent hexane (Hex), dichloromethane (DCM), and ethyl acetate (EtOAc) using a separatory funnel at room temperature. The solvents were removed using a rotary evaporator (Buchi R210; Switzerland, Flawil) at 35°C under vacuum. From the hexane fraction (GNSbM-F-Hex), 0.1330 g was obtained. The dichloromethane fraction (GNSbM-F-DCM) yielded 2.3697 g, the ethyl acetate fraction (GNSbM-F-EtOAc) yielded 10.5817 g, and the aqueous fraction (GNSbM-F-Aq) yielded 16.8156 g.

The hexane fraction (GNSbM-F-Hex) was submitted to column chromatography, with an extract/silica ratio of 1:200. The column was eluted according to a gradient of increasing polarity, from DCM:MeOH::H2O (67:28:5) to DCM:MeOH::H2O (55:40:5), obtaining a total of 12 fractions GNSbF-Hex (GNSbF-Hex-1–12). The GNSbF-Hex-3 fraction was purified by column chromatography, with an isocratic system of Hex:EtOAc (90:10) to get compound 1. The spectral properties of this compound, including 1H-NMR and 13C-NMR data, were identical to those previously described in the literature for the lupeol.

Lupeol (1) C30H50O crystals, m.p. 215–216°C. 1H-NMR (CDCl3, 400 MHz): δ (ppm); 4.68, 4.56 (2H, s, H-29a, 29b), 3.18 (1H, dd, J = 4.76 Hz, 11.2 Hz, H-3), 1.25, 1.02, 0.96, 0.94, 0.82, 0.78, 0.75, (each 3H, s, CH3×7). 13C-NMR (CDCl3, 100 MHz); δ (ppm); 38.2 (C-1), 25.3 (C-2), 79.2 (C-3), 38.7 (C-4), 55.4 (C-5), 18.5 (C-6), 34.4 (C-7), 40.9 (C-8), 50.6 (C-9), 37.3 (C-10), 21.1 (C-11), 27.6 (C-12), 39.0 (C-13), 42.9 (C-14), 27.6 (C-15), 57.5 (C-16), 42.9 (C-17), 48.5 (C-18), 48.1 (C-19), 151.1 (C-20), 29.9 (C-21), 40.2 (C-22), 28.1 (C-23), 15.5 (C-24), 16.3 (C-25), 16.2 (C-26), 14.7 (C-27), 18.5 (C-28), 109.5 (C-29), 19.5 (C-30).

The ethyl acetate fraction (GNSbMF-EtOAc) was purified by column chromatography, with an extract/silica ratio of 1:200. The column was eluted with DCM:MeOH::H2O 85:25:4 obtaining a total of 11 fractions GNSbMF-EtOAc (GNSbMF-EtOAc-1–11). The fraction GNSbMF-EtOAc-2 was also purified by column chromatography, with an isocratic system of EtOAc:MeOH 90:10 to get compound 2, identified as 3α-O-methyl ellagic acid 4-O-β-D-rhamnopyranoside.

3α-O-Methyl ellagic acid 4-O-β-D-rhamnopyranoside (2) C31H38O12, white amorphous powder, m.p. 248–250°C. 1H-NMR (CD3OD, 400 MHz): δ (ppm); 5.88 (1H, s, H-5), 8.14 (1H, s, H-5′), 6.53 (1H, d, J = 1.5 Hz, H-1′), 4.98 (1H, br, H-2′), 4.88 (1H, d, J = 3.4 Hz, H-3′), 4.85 (1H, d, J = 3.4 Hz, H-5″), 4.51 (1H, t, J = 9.4 Hz, H-4″), 4.28 (3H, s, 3′-OCH3), 1.76 (3H, d, J = 6.2 Hz, CH3-6″); 13C NMR: (CD3OD, 100 MHz) δ (ppm); 113.5 (C-1), 113.3 (C-1′), 135.9 (C-2), 143.2 (C-2′), 142.7 (C-3), 139.5 (C-3′), 146.7 (C-4), 152.8 (C-4′), 111.3 (C-5), 110.7 (C-5′), 114.3 (C-6), 113.4 (C-6′), 158.8 (C-7), 158.0 (C-7′), 100.7 (C-2′), 71.1 (C-3′), 72.8 (C-4′), 70.4 (C-5′), 17.2 (C-6″), 59.8 (3′-OMe).
The fraction GNSbMF-EtOAc-3 was purified by column chromatography, with an isocratic system of EtOAc: MeOH 85:15, to get compound 3, identified as 19-α-hydroxy-asiatic acid monogluconidade.

19-α-Hydroxy-asiatic acid monoglucoside (3) C_{36}H_{58}O_{11}, white amorphous powder, m.p. 220–221°C. 1H-NMR (DMSO-d6, 100 MHz): δ (ppm): 5.23 (1H, d, J = 7.9 Hz, H-1'), 4.29 (1H, s, H-2), δ: 4.15 (1H, m, H-5'); 4.28–4.57 (5H, m, H-2', H-3', H-4', H-6'); 3.82 (1H, d, J = 9.3 Hz, H-3), 5.23 (1H, m, H-12) 3.12 (1H, d, J = 14.3 Hz, H-18), 1.27, 1.08, 0.94, 0.65, 0.53 (each 3H, s, CH_{3}).

2.5. Morphological Analysis. Both cell lines were exposed to the antiproliferative extracts: RKO and SW613-B3. G. neuberthii extracts at 50 μg/mL of GNSbM extract or 0.3 μg/mL Dxo. Detached and attached cells were harvested and washed with PBS. Cell pellets were obtained and resuspended in 100 μL of PBS, fixed with absolute ethanol, and maintained at −20°C for 24 h. Cells were washed with PBS and incubated in the dark for 30 min at room temperature in the staining buffer (50 μg/mL PI, 0.1% sodium citrate, 0.1% Triton-X-100, and 100 μg/mL RNase A). Cells in the G_{1}, S, and G_{2}/M-phase were subsequently analyzed using a FACS Canto II flow cytometer (Becton Dickinson, USA). Acquired data were analyzed using DIVA and ModFit LT software (Becton Dickinson).

2.9. Cell Cycle Analysis. Cell cycle distribution was evaluated using propidium iodide (PI, P4170, Sigma Aldrich, USA) staining processed according to the previous protocol, Bailon-Moscoso et al. [11]. In summary, cells were seeded in 6-well plates at a density of 1 x 10^{5} cells in 2 mL of medium per well and incubated for 24 h. Cells were then treated for 48 h with 20, 30, and 50 μg/mL of GNSbM extract or 0.3 μg/mL Dxo. Detached and attached cells were harvested and washed with PBS. Cell pellets were obtained and resuspended in 100 μL of PBS, fixed with absolute ethanol, and maintained at −20°C for 24 h. Cells were washed with PBS and incubated in the dark for 30 min at room temperature in the staining buffer (50 μg/mL PI, 0.1% sodium citrate, 0.1% Triton-X-100, and 100 μg/mL RNase A). Cells in the G_{1}, S, and G_{2}/M-phase were subsequently analyzed using a FACS Canto II flow cytometer (Becton Dickinson, USA). Acquired data were analyzed using DIVA and ModFit LT software (Becton Dickinson).

2.10. Cloning Assay. To evaluate the clonogenic capacity, 2.5 x 10^{3} cells were seeded in duplicate in 6 cm diameter Petri dishes in 2 mL of medium and incubated for 24 h. Cells were then treated for 48 h with 20, 30, and 50 μg/mL of GNSbM extract or 0.3 μg/mL Dxo, washed with BPS, incubated with 2 mL of complete medium for 7 days, and then processed according to Guarnán-Ortiz et al. [12]. Colony-forming ability data were expressed as a percentage relative to control.

2.11. Western Blot Analysis. To determine the induced cell death pathway, 20, 30, and 50 μg/mL of GNSbM extract were exposed on both cell lines for 48 h. Additionally, as positive controls: cells were exposed for 10 min to UV radiation (Osram, G30T8, 30W Germicidal UV-C Lamp, 254 nm) for apoptosis induction [13] or for 1 h to PBS for starvation-induced autophagy [14], before being harvested. Apoptotic and autophagic proteins were analyzed through Western blot analysis. The methodology was applied according to Bailon-Moscoso et al. [15]. Briefly, separated proteins from a SDS-PAGE were transferred to a PVDF membrane (IPVH00010, Immobilon-P, 0.45 μm, EMD/Millipore, Billerica, Boston, MA, USA) and incubated with primary antibodies: p53 (sc-81168), Beclin-1 (sc-48341), SQSTM1/p62 (sc-48402) (Santa Cruz Biotechnology, USA), PARP (#9542), Bax (#2774), Bcl-2 (#15071) LC3A/B (#12741), and β-Tubulin (#2128) (Cell Signaling Technology, USA), as indicated by the manufacturer for immunoblotting. Secondary antibodies, anti-mouse IgG, HRP-linked (#7076, Cell Signaling Technology, USA), and anti-rabbit IgG, HRP-linked (#7074, Cell Signaling Technology, USA), were
subsequently used. Immunoreactive bands were visualized using an enhanced chemiluminescence Luminata Crescendo Western HRP Substrate or Luminata Forte Western HRP Substrate (Millipore-Merck, Germany).

2.12. Statistical Analysis. Statistical analyses were carried out in GraphPad Prism 4 (GraphPad Software, USA). All data were reported as the means ± SEM of three independently performed experiments, as detailed in each figure. The statistical significance was obtained with one-way analysis of variance (ANOVA) followed by the Dunnett posttest. A P < 0.05 was considered to be statistically significant comparing the control to the samples.

3. Results and Discussion

3.1. Preliminary Phytochemical Study of Extracts. The phytochemical screening tests on extracts revealed the presence or absence of the main secondary metabolites and other phytochemicals based on the presence or absence of expected color changes (Table 1). The _G. neuberthii_ fruit extracts in methanol (GNFM) are richer in secondary metabolites; the methanolic extracts of stem bark (GNSbM) contained alkaloids, flavonoids, tannins, quinones, and saponins. The highest percentage of yield was obtained from the methanol fraction of leaves GNLM (22.97%), followed by that of hexane fraction of fruit GNFH (19.6%), and methanol fraction of stem bark GNSbM (9.4%). The lowest percentage of yield was obtained from the hexane fraction of seed GNSH (0.17%). There was also variation in the physical appearance of the extracts (Supplementary Material, S1).

3.2. Cytotoxic Effect on Human Tumor Cell Lines. In order to analyze the antiproliferative effect of _G. neuberthii_ on colon cancer cells, all the extracts obtained were then evaluated. Table 2 describes the viability percentages using 50 μg/mL of the different _G. neuberthii_ extracts or Dxo 0.3 μg/mL on the human tumor cell lines after 48 h of treatment. Doxorubicin, an anthracycline antibiotic, has proved to induce cytotoxicity, cell cycle arrest, and apoptosis in a wide variety of tumor cell lines, including colon cancer cells; however, its clinical use is limited due its cardiotoxicity effect [16-18]. Cell growth viability was measured using the MTS assay, considering the control as 100% of viability. As expected, the SW613-B3 cell line (with p53 mutated) was more resistant to the treatment, in contrast to the RKO cell line. The extract GNSbM, whose extracting method was by maceration in methanol, demonstrated to be the most cytotoxic, with values under 20% of viability on both wt p53 RKO and SW613-B3 with mutated p53.

According to these results, the IC$_{50}$ was calculated from the GNSbM extract (Figure 1(a)), which was 28 and 31.8 μg/mL in SW613-B3 and RKO, respectively. As observed, after exposing to the IC$_{50}$ of GNSbM, that is approximately the same, both cell lines decrease in cell population, suggesting therefore that the effect observed is not dependent on p53 status.

The marked difference between these cell lines reveals that the SW613-B3 cells have the potential for recovery after treatment because of the results observed in colony-forming ability assay, where the cells were exposed to three different doses of GNSbM, 20, 30, and 50 μg/mL. As observed in Figure 1(b), colony-forming ability decreases in a dose-dependent manner in both cell lines; however, the SW613-B3 cell line was less affected after 20 μg/mL (90%) and IC$_{50}$ (80%), in contrast to the RKO cell line, which was more sensitive (60%). To note, SW613-B3 cell line has demonstrated to be resistant to different treatments. [12] Nevertheless, colony-forming ability decreases dramatically in both cell lines at the higher concentration (50 μg/mL). On the other hand, p53 has a well-known role in cell cycle progression, which after transactivation induces cell cycle arrest [19, 20]; however, the statistical analysis of cell cycle distribution revealed that no significant cell cycle changes were occurring after the treatment of both wt and mutated p53 cell lines, as observed in those treated with Dxo (Figure 1(c)).

3.3. No Apoptosis Was Detected after Treatment with GNSbM. To further explore the type of cell death induced by the plant extract, both apoptosis and autophagy pathways were analyzed by Western blotting. It is also well known that p53 is involved in multiple cell death pathways, such as apoptosis and autophagy. [21, 22] In the apoptotic pathway, p53 is upregulated to transactivate and phosphorylate the Bax protein, thereby inducing the activation of the intrinsic apoptosis; at the same time Bcl-2, an antiapoptotic protein, is downregulated in the presence of p53. [22] As observed in Figures 2(a) and 2(b), no overexpression of Bax protein was detected in the RKO cell line, despite the overexpression of p53 in a dose-dependent manner; furthermore, an increase of Bcl-2 expression was detected in the Western blot assay (Figure 2(b)), although this did not reach statistical significance relative to the control (Figure 2(a)).

Additionally, in the SW613-B3 cell line, the upregulation of p53 was not observed in any of the GNSbM doses used as expected (Figures 3(a) and 3(b)). In both cell lines, no proteolysis of PARP-1 protein, an apoptotic marker [23, 24], was detected after the GNSbM treatments (Figures 2 and 3), in contrast with the cleavage visible in cells exposed to UV radiation. [25]

3.4. GNSbM Extract Induces Autophagy. Autophagy pathway is activated after starvation conditions [26, 27], which induces the formation of autophagic vesicles or vacuoles called autophagosomes at the level of the cytoplasm. Therefore, the presence of vacuoles in both cell lines, observed in morphological analysis (Figure 1(a)), suggested the possible activation of the autophagic pathway [12, 28]; thus, biomarkers for autophagy were evaluated. In this pathway, Bcl-2 is associated with Beclin-1. Once Beclin-1 is released, the autophagic pathway is activated [29]; LC3-I is converted into its active form LC3-II to induce the phagophore formation in the nucleation phase to enclose the obsolete proteins and organelles, tagged with p62, for degradation [30-33]. Forced autophagy could generate severe damage
that ends in the death of the cell [27]. In Figures 2(b) and 3(b), an upregulation of Beclin-1 was observed in both cell lines in a dose-dependent manner, although these results did not reach statistical significance. Dissociation between Bcl-2 and Beclin-1 is necessary for the initiation of autophagy; therefore, the increase of Bcl-2 observed in the RKO cell line (Figure 2(b)) could be explained by this dissociation. Likewise, an increase in LC-3II was detected in both the RKO and SW613-B3 cell lines in a dose-dependent manner. k+he final step in autophagy is the degradation of charges in which p62 is involved (being also degraded). [32] As observed in both cell lines, p62 results in a decrease (Figures 2(a) and 3(a)), therefore elucidating the termination of the autophagic process and cell death pathway. Remarkably, it is possible that autophagy is playing a double role in the SW613-B3 cell line. It is well known that the autophagy mechanism is also active in tumor cell lines and has a role in survival [22, 34]; as observed in the clonogenic assay, the SW613-B3 cell line is able to recover after treatment at the IC_{50} dose (Figure 1(b)).

### 3.5. Isolation and Identification of Secondary Metabolites Isolated from GNSbM

Phytochemical investigation of methanolic extract from the stem bark of *G. neuberthii* (GNSbM) led to the isolation and identification of three compounds identified as: lupeol (1), 3′-O-methyl ellagic acid 4-O-β-D-rhamnopyranoside (2), and 19-α-hydroxy-asiatic acid monoglucoside (3), whose structures are shown in Figure 4. k+he spectral properties of these known compounds, including the 1HNMR and 13CNMR data, were identical to those previously described in the literature [9].

#### Compound 1

was crystals. Its 1H NMR spectra in CDCl_3 showed six singlets, corresponding to tertiary methyl groups, between 0.75 and 1.02 ppm and a singlet at 1.25 ppm, typical of a methyl group in an isopropenyl system. Two olefinic protons at 4.68 and 4.56 ppm are consistent with the methylene group of the same propylenic system. 13C NMR data showed the characteristic signals of C-3 at 79.2 ppm, C-20 at 151.1 ppm, and C-29 at 109.5 ppm. [9]

#### Compound 2

was a white amorphous powder. Its 1HNMR spectrum in deuterated pyridine showed two singlet signals at δ 8.58 and 8.14. A broad signal at 5.31 ppm suggested a structure with some hydroxyls groups, which was corroborated with the few multiplets signals typical for carbohydrate moieties between 4.53 and 5.04 ppm. Among these signals, one singlet at δ 4.28 integrating for 3H suggested a methoxy group in this compound. The broad band carbon NMR spectra showed 21 signals including 11 quaternary carbons indicating a very conjugated aromatic structure. From these signals, two were assigned to α, β-unsaturated lactone carbonyls carbons at 158.0 and 158.2 ppm. The

### Table 1: Phytochemical constituents of extract from the aerial parts of *G. neuberthii*.

| Test | Fruit | Seed | Stem bark | Leaves |
|------|-------|------|-----------|--------|
|      | Hex   | EtOAc| MeOH      | Hex    |
|      | GNFH  | GNFE | GNFM      | GNSH   |
| Proteins | − | − | + | − | + | − | − | − | − | + |
| Carbohydrates | − | − | + | − | − | − | − | − | − | − |
| Fats | +++ | ++ | + | ++ | − | ++ | − | + | − | − |
| Alkaloids | ++ | ++ | ++ | − | − | − | − | − | − | − |
| Terpenoids, steroids | − | − | + | − | − | − | − | − | − | − |
| Flavonoids | + | + | + | − | − | − | − | − | − | − |
| Saponins | − | − | + | − | − | − | − | − | − | − |
| Quinones | − | − | ++ | − | − | − | − | − | − | − |
| Tannins | − | − | + | + | − | − | + | − | − | + |

Hex = hexane extract, EtOAc = ethyl acetate extract, MeOH = methanol extract, +++ = very strong positive, ++ = strong positive, + = fair positive, − = absent.

### Table 2: Cell Growth Viability: Cell lines were treated for 48 h with 50 μg/mL of *G. neuberthii* extract.

| Part of the plant | Extract | RKO | SW613-B3 |
|------------------|---------|-----|----------|
| Fruit            | GNFH    | 96 ± 6.79 | NE |
|                  | GNFE    | NE | NE |
|                  | GNFM    | 72.2 ± 6.38 | NE |
| Seed             | GNSEa   | 81.0 ± 7.78 | NE |
|                  | GNSM    | 74.7 ± 4.36 | 95.0 ± 5.31 |
| Stem bark        | GNSbEa  | NE | NE |
|                  | GNSbM   | 15.3 ± 3.61 | 10.4 ± 2.24 |
| Leaves           | GNLM    | 91.5 ± 4.95 | NE |
| Control          | 0.3 μg/mL Dxo | 10.8 ± 4.02 | 48.2 ± 5.62 |

Control was considered as 100% of cell viability; three independent experiments in triplicate were performed. NE = no effect.
signals corresponding to the carbohydrate, analyzed together with the proton spectra, indicated that it was the β-D-glucopyranoside. All the data obtained by NMR were ascribable to a derivative of ellagic acid. The comparison of physical and spectroscopic data [35] indicated that the isolated compound 2 was 3′-O-methyl ellagic acid 4-O-β-D-
glucopyranoside. Compound 3 was a white amorphous powder. The NMR spectra were taken in deuterated DMSO, submitted to analyses. The $^1$H NMR showed the resonances of five quaternary methyl groups at 0.52, 0.54, 0.94, 1.08, and 1.26 ppm, together with a methyl doublet at $\delta$ 0.86. An olefinic proton in 5.14 ppm suggested an ursane-type
triterpene for the aglycone of this glycosylated compound. Few multiplets between 3.8 and 4.60 ppm were ascribable to a sugar moiety. The carbon NMR spectra analyzed with the help of DEPT experiment revealed for the aglycone triterpene 30 carbons, including six methyl groups, nine methylene groups, counting one oxygenated, seven methine groups with two oxygenated at 77.7 and 67.7 ppm, and one olefinic group at 138.4 ppm; eight quaternary carbons comprising the olefinic at 127.2 ppm, one oxygenated at 72.4 ppm, and the carboxyl ester group at 175.8 ppm. Comparative study of the resonances of the sugar indicated that it was the β-D-glucoside. The 2D experiments, COSY, HMQC, and HMBC were in agreement with the proposed structure. All the data were compared with the literature [36, 37] to finally consider this compound as 19-hydroxy-asiatic acid monoglucoside.

Regarding this, lupeol (1), 3′-O-methyl ellagic acid 4-O-β-D-rhamnopyranoside (2), and asiatic acid-β-D-glucoside (3) are reported here for the first time in this genus. For instance, lupeol has been widely reported for its anticancer effect against various cancer cells, such as oral cancer, pancreatic cancer, gallbladder cancer, prostate cancer, and colorectal cancer. [38–41] Also, asiatic acid is known to be cytotoxic to several tumor cell lines. However, asiatic acid-induced cell death was mainly apoptotic, demonstrated in colon cancer RKO cells [42, 43]. Hence, the presence of these phytochemicals in GNSbM might be synergistically responsible for the autophagy-inducing effect, as suggested by our results.

Figure 3: GNSbM extract induces autophagy on SW613-B3 cell line. Cells were exposed to GNSbM or UV radiation or starvation in PBS. Total protein was separated in a SDS-PAGE followed by Western blot analysis with indicated antibodies against p53, apoptotic and autophagy biomarkers, and tubulin as a loading control. (a) Quantification of the level of p53 expression and the apoptotic and autophagy biomarkers. Data represented the mean ± SEM (n = 6) of three independent experiments. k+_he tests for significance were limited to the ANOVA-Dunnet posttest: ∗P < 0.01, ∗∗P < 0.001, ∗∗∗P < 0.0001 vs. control. (b) Western blot pictures demonstrating the effect observed in Sw613-B3 cell line. No p53 expression was observed as expected although autophagy activity was detected after treatment.
4. Conclusions

In summary, different extracts of G. neuberthii were evaluated; the one with the greatest cytotoxic effect was the so-called GNSbM, which considerably reduced cell viability in both RKO cell line, with wt p53, and SW613-B3, with mutated p53, in a dose-dependent manner after 48 h of exposure. In addition, activation of the apoptotic route was discarded and the evidence of autophagic activity was detected. Finally, three compounds were identified in this extract: lupeol, 3'-O-methyl ellagic acid 4-O-β-D-rhamnopyranoside, and asiatic acid-β-D-glucoside, which have been shown to have antitumor effects.

Data Availability

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

Conflicts of Interest

The authors declare that there are no conflicts of interest regarding the publication of this paper.

Authors’ Contributions

Luis M. Guaman-Ortiz and Juan C. Romero-Benavides contributed equally to this work.

Acknowledgments

This work was supported by International Centre for Genetic Engineering and Biotechnology (ICGEB), Trieste, Italy (ECU1601 EC) and Universidad Técnica Particular de Loja (UTPL), Loja, Ecuador (PROY_CCSAL_1266). The authors want to thank Dr. A. Ivana Scovassi for her support in the observations and analysis of experimental results and reviewing the English version.

Supplementary Materials

Different parts of the G. neuberthii were macerated for 72 h in a light-free environment with three dissolvents separately: hexano, ethyl acetate, and methanol. The extracts were concentrated at 50 mbar and 37°C on a rotary evaporator. The appearance and weight of the extracts were recorded, and the yield was calculated. (Supplementary Materials)

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