Phytochemical analysis and in vitro anti-proliferative activity of *Viscum album* ethanolic extracts

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**Abstract**

**Background:** *Viscum album* L. (Santalaceae), commonly known as mistletoe, is a hemiparasitic plant traditionally used in complementary cancer treatment. Its antitumor potential is mostly attributed to the presence of aqueous soluble metabolites; however, the use of ethanol as solvent also permits the extraction of pharmacological compounds with antitumor potential. The clinical efficacy of mistletoe therapy inspired the present work, which focuses on ethanolic extracts (*V. album* “mother tinctures”, MT) prepared from different host trees.

**Methods:** Samples from three European subspecies (*album*, *austriacum*, and *abietis*) were harvested, and five different *V. album*-MT strains were prepared. The following phytochemical analyses were performed: thin layer chromatography (TLC), high-performance liquid chromatography (HPLC) and liquid chromatography-high resolution mass spectrometry (LC-HRMS). The proliferation assay was performed with WST-1 after incubation of tumor (Yoshida and Molt-4) and fibroblast cell lines (NIH/3 T3) with different MT concentrations (0.5 to 0.05% v/v). The cell death mechanism was investigated by flow cytometry (FACS) using Annexin V-7AAD.

**Results:** Chemical analyses of MT showed the presence of phenolic acids, flavonoids and lignans. The MT flavonoid and viscotoxin contents (mg/g fresh weight) were highest in *Quercus robur* (9.67 ± 0.85 mg/g) and *Malus domestica* (3.95 ± 0.58 mg/mg), respectively. The viscotoxin isofrom proportions (% total) were also different among the VA subspecies with a higher content of A3 in *V. album* growing on *Abies alba* (60.57 ± 2.13). The phytochemical compounds as well as the viscotoxin contents are probably related to the antitumor effects of MT. The cell death mechanisms evaluated by colorimetric and FACS methodologies involved necrotic damage, which was host tree-, time- and dose-dependent, with different selectivity to tumor cells. Mother tincture from *V. album* ssp. *abietis* was the most effective at inducing in vitro cellular effects, even when incubated at the smallest concentration tested, probably because of the higher content of VT A3.

**Conclusion:** Our results indicate the promising antitumor potential of *Viscum album* ethanolic extracts and the importance of botanical and phytochemical characterization for in vitro anti-proliferative effects.

**Keywords:** Mistletoe, *Viscum album* L., Phytochemistry, Anti-proliferative activity, Cytotoxicity

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Background

Viscum album L. (Santalaceae), also known as European mistletoe, is a hemiparasitic shrub that is differentiated into 3 main subspecies growing on different host trees, i.e., V. album ssp. album, deciduous trees; V. album ssp. abietis, fir, and V. album ssp. austriacum, mainly on pine [1].

Most investigations into V. album are based on aqueous extracts [2–6]. The active compounds that have been identified are proteins commonly classified as visotoxins (VT) and mistletoe lectins [7, 8]. Phytochemical investigations of V. album also revealed the presence of other important pharmacological compounds, such as phenolic acids, phenylpropanoids, flavonoids, triterpenes, phytosterols, oligopeptides and polysaccharides [9, 10].

The European Medicines Agency reported the traditional use of different V. album ethanolic extracts to treat cardiovascular disease [11]. Additionally, Porruthukaren et al. [12] described a reduction in blood pressure after V. album ethanolic use for 12 weeks.

Nevertheless, the antitumor effects of ethanolic V. album extracts in biological systems have been described. In vivo studies have shown the anticancer activity of alcoholic and glycerine V. album extracts by stimulating immune mechanisms and inhibiting tumor cell proliferation [13]. The simultaneous application of V. album ethanolic extract and doxorubicin increases the toxicity in Ehrlich tumor cells, opening up the possibility of exploring a tentative synergy between single chemical compounds and complex herbal extracts [14, 15]. In addition, the in vitro research performed with these extracts highlighted the apoptotic mechanisms and cell growth reduction [16, 17] involved in melanoma, leukemia [16], and human cervix adenocarcinoma tumor [17] death. Panossian et al. [18] suggested that phenylpropanoids detected in V. album ethanolic extracts had antitumor activity through the inhibition of protein kinase C.

Furthermore, a recent study performed with V. album ethanolic extracts showed tumor cell cycle arrest and apoptotic death in vitro models. Chemical analysis of these identified compounds such as caffeic acid, chlorogenic acid, sakuranetin, isosakuranetin, syringenin 4-O-glucoside, syringenin 4-O-apiosyl-glucoside, alangilignoside C and ligalumioside A [16].

It is widely known that many metabolites isolated from European mistletoe are not produced by the plant itself but are due to host tree metabolism [19], supporting the importance of understanding the subspecies of European mistletoe and the influence of the host tree. This work shows, for the first time, the in vitro anti-proliferative effects and the chemical composition of ethanolic extracts produced from different V. album host trees.

Methods

Plant growth and harvest

The green, unripe berries, leaves and stems of three European subspecies of V. album L. were harvested in July 2016 in natural habitats in Switzerland (Fig. 1a-g). The following V. album subspecies were collected from five different host trees: V. album ssp. album growing on Malus domestica (VAM; Fig. 1c), Quercus robur (VAQ; Fig. 1d) and Ulmus carpinifolia (VAU; Fig. 1e); V. album ssp. abietis growing on Abies alba (VAA; Fig. 1f) and V. album ssp. austriacum growing on Pinus sylvestris (VAP; Fig. 1g). From each host tree, at least five bushes of the same age containing the same parts of the plant were submitted to solvent extraction as follows: first and second youngest leaves, berries, and first and second youngest stems (Fig. 1a-b) were harvested five times. The samples were always collected in the morning (from 8:00 to 11:30 am) and immediately transported to perform ethanolic extraction within 4 to 6 h.

All plants were identified by Dr. Marcelo Guerra Santes (Universidade Estadual do Rio de Janeiro). Voucher specimens (C.H. Quaresma 18.328, C.H. Quaresma 18.329, C.H. Quaresma 18.332, C.H. Quaresma 18.327 and C.H. Quaresma 18.331) were deposited at the Herbarium of the Faculdade de Formação de Professores, Universidade Estadual do Rio de Janeiro, Brazil.

Preparation of V. album mother tinctures

All solvents and reagents exhibited analytical purity quality. The fresh material (5 g) was fragmented into segments smaller than 5 cm long and dried in an oven at 105 °C for 2 h, following the Brazilian Homeopathic Pharmacopeia [20] and the French Pharmacopoeia [21]. Once the percentage of solid residue of each fresh plant had been established, the total volume of mother tincture (MT), as well as the volume and the concentration of ethanol used in the maceration process, was determined. Next, the maceration extractive process was conducted over a period of 3 weeks at room temperature in 80% w/w ethanol. To increase the efficiency of the extraction process, all V. album ethanolic solutions were shaken by hand for 60 s twice a day. After 3 weeks, the macerates were filtered and kept at 20 ± 5 °C. The final MT concentrations were 40 to 50% v/v [21].

Thin layer chromatography

Thin layer chromatography (TLC) was performed as specified in the monograph of V. album in the French Pharmacopoeia [21]. Samples (10–20 µL/band) were applied as 10 mm bands with a 10 mm track distance onto the plate by a capillary 10 mm from the lower edge of the plate. TLC separations were achieved in a chamber (210 × 100 mm) saturated with distilled water:methanol:glacial acetic acid:dichloromethane at 2:3:8:15 (v/v/v/v/v).
as a mobile phase. The bands were visualized under UV light (365 nm) before and after spraying with NP/PEG as the revealing solution.

**Determination of total flavonoid content**

The concentration of total flavonoids as rutin equivalents was determined spectrophotometrically in the UV region (360 nm) in comparison with the standard curve of rutin absorption, adapted from Rolim et al. [22]. The concentration range of the standard curve comprised the following concentrations of rutin: 5.0, 10.0, 15.0, 20.0, 25.0 and 30.0 μg/mL. The absorptions of rutin concentration series were plotted to provide a linear calibration curve with $r^2$ close to 1. A mixture of 95% ethanol and 0.02 M acetic acid (99:1) was used as the solvent in all solution preparations. The readings were performed in triplicate.

**HPLC viscotoxin analysis**

Before HPLC analysis, all mother tincture samples were purified by solid phase extraction (Bakerbond, carboxylic acid, wide pore SPE column) to separate the VT from mother tincture impurities. For this, the SPE column was previously washed with methanol and water and subsequently equilibrated with 5 mL of a 200 mM ammonium solution. Aliquots from 0.5 to 2 mL of each mother tincture were added to the column and the pH was adjusted to 7.0–7.5. The samples were eluted...
under vacuum, and 3 mL of water was used to rinse the samples. Finally, the samples were eluted with 5 mL of 0.4 M acetic acid and the HPLC methodology described by Schaller et al. [7] was used for VT quantification. This involved injecting crude acid extracts into the column without further clean up. Column Nucleosil C18 AB, 125 × 4 mm with guard column; eluent A 0.1% TFA in water; eluent B: 0.1% TFA in acetonitrile/water = 60/40. The gradient was as follows: 0–9.0 min, 38% B to 42% B; 9.0–9.5 min, 42% B to 50% B; 9.5–17.0 min, 50% B to 54% B; 17.0–18.0 min, 54% B to 70% B; 18.0–19.0 min 70% B; 19.0–19.2 min 70% B to 38% B; 19.2–21.0 min 38% B; flow rate: 1 mL/min; detection, UV at 210 nm. The VT eluted in the order B, A1, A2, A3 and the total visco-toxin content (μg/mL) was calculated as the sum of each isoform.

**LC-HRMS analysis**

Samples were prepared as described in the *V. album* monograph with modifications [21]. UHPLC Dionex Ultimate 3000 coupled to a Q Exactive Plus Orbitrap mass spectrometry system (Thermo Fisher Scientific, Germany), equipped with an electrospray ionization (ESI) source operating in negative ion mode at a voltage of 2.9 kV was employed for the chemical analysis. Separations were performed on a reversed-phase column (Thermo Syncronis C18 50 mm × 2.1 mm; 1.7 μm). Water-formic acid 0.1% v/v (A) and acetonitrile (B) were used as mobile phases as follows: (0–15 min, 2% B; 15–16.2 min, 20% B; 16.2–17 min, 100% B; 17–18 min, 100–2% B; 18–19.5 min 2% B). The flow rate was 0.4 mL/min and the injection volume was 5 μL. The mass spectra were acquired in full scan mode at a resolution of 70,000 over a *m/z* range of 60–780. Chromatograms were aligned using the XCMS online platform, and compound annotation was performed by comparing the exact mass and MS/MS spectra of compounds (Supplementary information) with those available in different databases (European MassBank, MassBank of North America – Mona, SciFinder Scholar, NIST MSMS 2014) and in the literature for *V. album*. A 5 ppm tolerance error between the theoretical and experimental mass values was considered for compound annotation.

**Cell lines and culture conditions**

The following cell lines were used: MOLT-4 (human acute lymphoblastic leukemia cell line), Yoshida (mouse sarcoma cell line) and non-tumor NIH/3 T3 (mouse embryonic fibroblasts cell line), obtained from ATCC, Rockville, MD, USA (Yoshida and MOLT-4) and from the German Collection of Microorganisms and Cell Cultures GmbH, Braunschweig, Germany (NIH/3 T3). All cells were cultured in RPMI-1640 medium supplemented with 5% heat-inactivated fetal calf serum (FCS), 2 mM L-glutamine, and 1% penicillin-streptomycin in a humidified atmosphere with 5% CO₂ at 37 °C. Cell lines were maintained in exponential growth, and cells from sub-confluent monolayers (Yoshida and NIH/3 T3) were harvested by trypsin-EDTA to carry out the experiments.

**Cell viability assay**

Cell viability was evaluated by the WST-1 colorimetric methodology. Briefly, 90 μL of each cellular suspension containing 5 × 10⁴ cells/mL was pre-cultured in 96-well plates. After 24 h, 90 μL of *V. album* mother tinctures, pre-diluted in cellular culture medium, were added at concentrations varying from 0.05 to 0.5% v/v. Cellular viability rates were measured after incubation for 4 and 24 h by the addition of 20 μL of WST-1 to each well. The absorption at 450 nm and 650 nm against a background control was measured after 3 h of incubation at 37 °C in the dark in a multiwell plate reader. Since ethanol was used as the solvent for the extraction of tinctures, its effect on cell viability was also evaluated using the same MT concentrations. The percentage of viable cells was calculated in relation to control cells (untreated and treated with ethanol solvent) using mean values from at least three independent experiments, and the calculation was performed in triplicate. The IC50 was calculated using GraphPad 5 Software.

**Determination of apoptosis/necrosis by flow cytometry**

Molt-4 and Yoshida cells were seeded on 6-well plates at a concentration of 1 × 10⁵ cells/well. After 24 h, cells were treated with the following MT concentrations (% v/v) for 4 and 24 h: VAM (0.15 and 0.5), VAQ (0.15 and 0.5), VAU (0.35 and 0.5), VAA (0.05 and 0.5), and VAP (0.35 and 0.5). After treatments, the cellular suspension (Molt-4) and supernatant (Yoshida) were collected. Trypsin was added to the supernatants of the Yoshida cells. Then, apoptosis/necrosis was measured using the Annexin V-FITC Apoptosis Detection Kit I (BD Biosciences Pharmingen). Apoptotic cells: Annexin V-FITC Apoptosis Detection Kit I (BD Biosciences Pharmingen). Apoptotic cells: Annexin V-FITC positive and 7-AAD negative. Late apoptotic/necrotic cells: Annexin V-FITC positive and 7-AAD positive. Values are given as the percentage of total cells in relation to the ethanol control.

**Statistical analysis**

In vitro experiments were performed at least three different times, and results were analyzed by ANOVA with Dunnett’s post hoc test using GraphPad 5. *P* values < 0.05 were considered statistically significant.
Results and discussion

Chemical analysis of V. album ethanolic extracts
TLC plates of ethanolic extracts showed orange-yellowish bands at 365 nm UV light after spraying with NP/PEG reagent, and this result is typical of flavonoid compounds. Each MT exhibited one fluorescent blue spot with a Rf (retention factor) value similar to that of the chlorogenic acid standard (Rf 0.60). According to ANSM [21], chlorogenic acid is a marker of V. album species, and its identification is important to assure the quality of the vegetal material. Łuczkiwicz et al. [23] also identified chlorogenic acid in alcoholic extracts of V. album, corroborating the results found in this work.

Determination of the total flavonoid content of the mother tincture
Table 1 shows the flavonoid concentration expressed as mg/g of plant fresh weight (mg/g fw) after ethanolic extraction. The highest concentration was detected in V. album growing on Quercus robur (9.67) and Malus domestica (6.30). Pietrzak et al. [24] observed an increase in the flavonoid content of V. album ssp. abietis extracted by a mixture of polar organic solvents with water. Additionally, Pietrzak et al. [25] determined that the flavonoid content in methanol extracts from V. album growing on different host trees ranged from 0.270 to 0.428 mg/g. The host species and the organ harvested influenced the chemical composition of mistletoe [8].

HPLC viscotoxin analysis
Quantitative analysis (Table 1) of VT isoforms A1, A2, A3, B and 1-PS in ethanolic extracts showed that V. album ssp. abietis contains predominantly viscotoxin A3, whereas V. album ssp. austriacum contains viscotoxins A2 and 1-PS, which were not detected in V. album ssp. album. The three European subspecies of V. album could be distinguished on the basis of their VT composition, and this result is in accordance with literature data [26, 27]. However, the total content of VT is greater in aqueous preparations than in hydroalcoholic preparations when the viscotoxin proportions are compared for the same V. album ssp. [27].

LC-HRMS analysis
This analysis was performed to obtain an overview and compare the chemical compounds of the different V. album MTs prepared. The major peaks present in the LC-HRMS chromatograms were tentatively identified, and this compound identification may help in the standardization of the V. album extracts used in the in vitro analysis. Regarding the chemical complexity, a total of seven compounds (1–7, Fig. 2) were putatively annotated based on their highly accurate m/z values and MS/MS fragmentation spectra (see Supplementary material Fig. S1–S6). These peaks presented deprotonated molecular ions at m/z: 191.05585 [M-H]− (C7H11O6), 353.08908 [M-H]− (C16H17O9), 353.08908 [M-H]− (C16H17O9), 625.14038 [M-H]− (C27H29O17), 565.15607 [M-H]− (C26H29O14), and 581.22496 [M-H]− (C28H37O13) (Table 2).

Compound 1 presented an ion at m/z 191.05585 [M-H]−, similar to quinic acid C7H12O6 (~1.36 ppm error) [28]. Quinic acid has already been described in the alcoholic extract of V. schimperi [29].

Compounds 2, 3 and 4 were chlorogenic acid isomers and showed similar deprotonated molecular ions at m/z 353.08908 [M-H]− in the following elution order: 3-O-caffeoylquinic acid, 5-O-caffeoylquinic acid and 4-O-caffeoylquinic acid, respectively (C16H18O9, 3.62 ppm error). This order is in accordance with Zhang et al. [30], who also used a reverse-phase C18 column. Chlorogenic acid has already been reported in the ethanol extract of V. album [16, 23] and is important to assure the quality of the vegetal material.

Compound 5 presented an ion at m/z 625.14038 [M-H]− and was tentatively identified as the flavonol quercetin-di-hexoside. The MS/MS spectrum of this compound showed a characteristic loss of two hexose units (M-162 Da).

| V. album Subspecies | Mother tincture/Host tree | Viscotoxin content (% total) | Total Viscotoxin (mg/g fw) | Total Flavonoid (mg/g fw ± sd) |
|-------------------|---------------------------|----------------------------|--------------------------|-------------------------------|
|                   | A1 | A2 | A3 | B | 1-PS |                   |                       |
| abietis           | n.d | n.d | 60.57 ± 2.13 | 0.58 ± 0.35 | n.d | 2.48 ± 0.80 | 5.25 ± (1.70) |
| austriacum        | n.d | 60.82 ± 18.23 | 24.17 ± 15.00 | n.d | 33.09 ± 20.08 | 0.34 ± 0.25 | 4.55 ± (1.44) |
| Malus domestica   | 7.76 ± 3.71 | 49.50 ± 2.68 | 39.19 ± 1.12 | n.d | n.d | 3.95 ± 0.58 | 6.30 ± (1.77) |
| album             | 11.57 ± 3.10 | 63.25 ± 6.45 | 23.66 ± 5.84 | n.d | n.d | 3.08 ± 0.77 | 9.67 ± (0.85) |
| Ulmus carpinifolia| 13.64 ± 4.29 | 46.09 ± 8.97 | 22.36 ± 1.94 | 17.91 ± 10.43 | n.d | 2.42 ± 1.49 | 4.67 ± (1.37) |

Table 1 Total viscotoxin and flavonoid contents (mg/g fw) and proportions of the viscotoxin isoforms A1, A2, A3, B, and 1-PS in the Viscum album mother tincture samples, n.d: not detectable.
Compound 6 showed an ion at \( m/z \) 565.15607 \([\text{M-H}]^-\) and a fragment ion at \( m/z \) 271 \([\text{M-H}]^-\), which correspond to an aglycone moiety of a flavonoid structure, and two sugar unit losses: hexose (M-162 Da) and pentose (M-132 Da). Thus, compound 6 was identified as pinobanksin or naringenin pentoside-hexoside \((C_{26}H_{30}O_{14}, 0.36 \text{ ppm error})\) [31–33]. This compound showed a higher intensity signal in VAA compared to the other ethanolic extracts analyzed.

Compound 7 presented a deprotonated molecular ion peak at \( m/z \) 581.22496 \([\text{M-H}]^-\) with a fragment ion at \( m/z \) 419.17270, suggesting the loss of hexose. These compounds,

**Table 2** Chemical compounds putatively identified in *Viscum album* by LC-HRMS/MS in negative mode ionization

| Observed \( m/z \) [\text{[M-H}]^-] | Theoretical \( m/z \) | Error (ppm) | Chemical name/ molecular formula | MS/MS fragment ions | Chemical class |
|----------------------------------|----------------------|-------------|----------------------------------|---------------------|---------------|
| 1 191.05585                      | 191.05611            | 1.36        | Quinic acid \((C_7H_4O_3)\)      | 191.05558 111.00785 87.00770 | Phenolic acid |
| 2 353.08908                      | 353.08780            | 3.62        | 3-O-coffeoylquinic acid \((C_{9}H_{8}O_5)\) | 191.05595 179.03467 135.04451 | Phenolic acid |
| 3 353.08908                      | 353.08780            | 3.62        | 5-O-coffeoylquinic acid \((C_{9}H_{8}O_5)\) | 191.05595 179.03467 135.04451 | Phenolic acid |
| 4 353.08908                      | 353.08780            | 3.62        | 4-O-coffeoylquinic acid \((C_{9}H_{8}O_5)\) | 191.05595 179.03467 135.04451 | Phenolic acid |
| 5 625.14038                      | 625.14102            | - 1.02      | Quercetin-di-hexoside \((C_{21}H_{30}O_{15})\) | 463.08899 301.03564 | Flavonoid     |
| 6 565.15607                      | 565.15627            | - 0.36      | Pinobanksin or naringenin pentoside-hexoside \((C_{20}H_{30}O_{14})\) | 271.06149 | Flavonoid     |
| 7 581.22496                      | 581.22396            | 1.72        | Lyoniresinosil-hexoside \((C_{20}H_{30}O_{14})\) | 419.17270 401.16071 | Lignan        |
already described in the genus *Viscum* [34], were assigned as lyoniresinol-hexoside (C_{28}H_{38}O_{13}), with a −1.72 ppm error [35].

**Cell viability assay**

The toxicity of the alcoholic vehicle was first evaluated by WST assay in Molt-4 and Yoshida tumor cells, and no cellular toxicity was observed (Fig. 3a-b). Therefore, the next tests with MT were performed with vehicle concentrations between 0.05 and 0.5% v/v.

Molt-4 (Fig. 3a) and Yoshida cells (Fig. 3b) exhibited approximately the same sensitivity to the tinctures, with MT from *Abies alba* (VAA) being the most effective, with IC50 values of 0.07 ± 0.01% v/v and 0.05 ± 0.03% v/v for Molt-4 and Yoshida, respectively. The non-tumor cells (3T3) were less sensitive with IC50 values of 1.60 ± 0.48% v/v after incubation with VAA. VAP did not exhibit toxicity in Molt-4 cells at any concentration tested (Fig. 3a) and exhibited only slight toxicity in Yoshida cells (p < 0.05; concentration range from 0.35–0.50, Fig. 3b).

The anti-proliferative assay highlighted the antitumor potential of MT within 4 h of incubation (Fig. 4a-b), in which a significant reduction in viability of approximately 90% was detected (0.5% v/v VAA, VAM, VAQ; p < 0.0001).

Flow cytometry was used to characterize the mechanism of cell death induced by *V. album* MT after 4 and 24 h of incubation (Fig. 5a-b). The number of necrotic cells increased proportionally to the MT concentration and to the incubation time, with a very similar profile for both cell lines, except in response to VAP. In Molt-4 cells, 0.5% v/v VAA, VAM and VAQ induced 61, 31 and 21% necrotic cell death, respectively, after 4 h of incubation (Fig. 5a top). Similar percentages of necrosis were observed in Yoshida cells (66, 47 and 35%), respectively, with the same MT concentrations. The necrotic effects were more evident after 24 h, supporting the necrotic potential of *Abietis, Malus* and *Quercus* MT (Fig. 5b bottom).

The present data suggest that the antitumor activity of MT from *V. album* ssp. *abietis* may be related to its higher content of VT A3 in addition to the presence of small molecules, as discussed below. Indeed, Schaller et al. [26] demonstrated that the viscotoxin A3 of *V. album* aqueous extract was more cytotoxic than other VT isoforms when the Yoshida sarcoma cell line was evaluated [26]. Additionally, Coulon et al.
attributed the highest antitumor activity of viscotoxin A3 to its physicochemical characteristics, in which the hydrophobic residues and the net liquid charge were able to increase the cell membrane interaction.

In addition, in vitro studies revealed that methanol extracts of mistletoe berries and ethanolic tinctures from whole *V. album* decreased the proliferation of colon cancer cells and viability of the murine melanoma lineage in a dose-dependent manner, respectively [16, 25]. These cellular alterations were attributed to modifications of mitochondrial activity and the cell cycle, among other cellular damage. Both studies showed that the polyphenolic composition of these *V. album* extracts was involved in the antitumor activity.

Regarding the seven compounds tentatively identified in this study, compound 1 was able to promote cytotoxicity in squamous cell carcinoma 4 (SCC-4). Compounds 2, 3 and 4 were chlorogenic acid isomers formed by quinic and cinnamic acid esterification [37]. Recently, Pan et al. [40] demonstrated that a type of quercetin-di-hexoside (quercetin 3,4′-di-O-glucoside) was an effective inhibitor of the growth of HepG2, PC3 and HT29 cells, corroborating the antitumor potential of *V. album* ethanolic extracts.

Pinobanksin or naringenin pentoside-hexoside (compound 6) presented a higher peak intensity in *V. album* extracts prepared with *V. album* ssp. *abietis* when compared to the other ethanolic extracts. The antitumor potential of these flavanones was previously described in tumor cell lines [41, 42]. In addition, the metabolomics analyses performed by our group with 50 different *V. album* samples harvested in winter and summer seasons from the same habitat in Switzerland also confirmed the importance of compound 6 in subspecies sample differentiation (*data not shown*).

Compound 7 was identified as lyoniresinol-hexoside, and its aglycone has already been described in *V. album* ssp. *coloratum* [35]. This lignan had cytotoxic properties against B16F10 cells after 48 h of treatment [43]. In addition, Baek et al. [44] demonstrated the ability of (−)-9′-O-(α-L-Rhamnopyranosyl) lyoniresinol to suppress A2780 human ovarian carcinoma cell proliferation in a dose-dependent manner.

**Conclusion**

The present study shows the antitumor potential of *V. album* tinctures in in vitro models. The cell death mechanism involved necrotic effects depending on the influence of the host tree, time and dose. Mother tincture from *V. album* ssp. *abietis* growing on *Abies alba* was the most effective, probably because of the higher content of VT A3, while MT from *V. album* ssp. austriacum growing on *Pinus sylvestris* exhibited only a slight

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**Fig. 4** WST-1 assay after 4 h of incubation with different concentrations of five *Viscum album* mother tinctures (VAM, *V. album* ssp. *album* growing on *Malus domestica*; VAQ, *V. album* ssp. *album* growing on *Quercus robur*; VAU, *V. album* ssp. *album* growing on *Ulmus carpinifolia*; VAA, *V. album* ssp. *abietis* growing on *Abies alba*; and VAP, *V. album* ssp. austriacum growing on *Pinus sylvestris*). a Molt-4 and b Yoshida cell lines. The results are presented as the mean ± SD from three independent experiments in relation to control cells. ***p < 0.0001, obtained with one-way ANOVA with Dunnet’s post hoc test. Legend symbols: ■3OH, cells incubated with ethanol solvent; □Control, untreated cells; ▪*V. album* ssp. *album* growing on *Malus domestica* (VAM), *Quercus robur* (VAQ) and *Ulmus carpinifolia* (VAU); ▼*V. album* ssp. *abietis* growing on *Abies alba* (VAA); and ◇*V. album* ssp. austriacum growing on *Pinus sylvestris* (VAP).
antitumor effect. Additionally, tumor cells were more sensitive than normal fibroblasts, suggesting that *V. album* MT has promising antitumor potential. The small molecules identified in MT were quinic acid, three isomers of chlorogenic acid, quercetin-di-hexoside, pinobankasin or naringenin pentoside-hexoside and lyoniresinol-hexoside. Further studies using in vitro and in vivo models in addition to stability assessments, should be performed to stimulate the development of new pharmaceutical formulations containing *V. album* ethanolic extracts.

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s12906-020-02987-4.

**Additional file 1.**

**Abbreviations**

EDTA: Ethylenediaminetetraacetic acid; ESI: Electrospray ionisation; FCS: Fetal calf serum; HPLC: High-performance liquid chromatography; HRMS: High-resolution mass spectrometry; IC50: Half maximal inhibitory concentration; LC-HRMS: Liquid chromatography-high resolution mass spectrometry; MS: Mass spectrometry; MT: Mother tincture; NP/PEG: Diphenylboriloxyethilamine/polyetileneglicol; PDA: Photodiode array detector; RF: Retention factor; TIC: Total ion current; TLC: Thin layer chromatography; VAA: *V. album* ssp. abietis growing on *Abies alba*; VAM: *V. album* ssp. album growing on *Malus domestica*; VAQ: *V. album* ssp. album growing on *Quercus robur*; VAU: *V. album* ssp. austriacum growing on *Ulmus carpinifolia*; VAA, *V. album* ssp. abietis growing on *Abies alba*; VAP, *V. album* ssp. austriacum growing on *Pinus sylvestris*.

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**Authors’ contributions**

CH and MNOM were responsible for the general study concept, conducting experimental analysis, interpreting and discussing of results, and drafting the
manuscript. APO and RG analyzed and interpreted the chemical experiments. JVCB performed and described the flavonoid experiments. MAMC interpreted and discussed the results. MG and CH harvested the Viscum album samples and were also involved with ethanolic extract preparation. HR and GS contributed to the study concept. CDT was responsible for the viscostoxin HPLC analysis. UW contributed to the in vitro cell viability experiments and data interpretation. KU and SB supported the study concept and data analysis. CH and SB were the study coordinators. All authors have read and approved the final manuscript.

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Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

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Not applicable.

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