Article

Extract of *Herba Anthrisci cerefolii*: Chemical Profiling and Insights into Its Anti-Glioblastoma and Antimicrobial Mechanism of Actions

Dejan Stojković 1, Danijela Drakulić 2, Marija Schwirtlich 2, Nemanja Rajčević 3, Milena Stevanović 2,3,4, Marina D. Soković 1 and Uroš Gašić 1,

1 Department of Plant Physiology, Institute for Biological Research “Siniša Stanković”—National Institute of Republic of Serbia, University of Belgrade, Bulevar despot Stefan 142, 11000 Belgrade, Serbia; dejanbio@ibiss.bg.ac.rs (D.S.); mris@ibiss.bg.ac.rs (M.D.S.)
2 Institute of Molecular Genetics and Genetic Engineering, University of Belgrade, 11042 Belgrade, Serbia; danijeladrakulic@imgge.bg.ac.rs (D.D.); schwirtlich@imgge.bg.ac.rs (M.S.); milenastevanovic@imgge.bg.ac.rs (M.S.)
3 Faculty of Biology, University of Belgrade, 11000 Belgrade, Serbia; nemanja@bio.bg.ac.rs
4 Serbian Academy of Sciences and Arts, 11001 Belgrade, Serbia
* Correspondence: uros.gasic@ibiss.bg.ac.rs

Abstract: *Anthriscus cerefolium* (L.) Hoffm. is a plant traditionally used around the globe since antiquity. Although widely used in many traditional medicines in different cultures, from the scientific point of view it is poorly investigated. Glioblastoma, a tumor type with poor prognosis, is the most common and lethal brain tumor in adults. Current therapeutic strategies for glioblastoma include surgery, radiation and chemotherapy. On the other hand, it has been revealed that patients with cancers are highly susceptible to microbial infections due to the invasive nature of cancer treatment approaches. This study was designed to investigate the chemical profile of *herba Anthrisci cerefolii* methanolic extract by applying UHPLC-LTQ Orbitrap MS4 analysis and to analyze its anti-glioblastoma and antimicrobial activities. This study revealed that methanolic extract of *herba Anthrisci cerefolii* contained phenolic acids and flavonoids, with 32 compounds being identified. Anti-glioblastoma activity was investigated in vitro using A172 glioblastoma cell line. The cytotoxic effects of the extract on A172 cells were compared to the same effect on primary human gingival fibroblast (HGF-1) cells. Decreased rate of proliferation and changes in cell morphology were detected upon treatment of A172 cells with the extract. The antimicrobial activity of extract was tested against *Staphylococcus aureus* and *Candida* species. The extract was active against the tested bacterium and yeasts, inhibiting free floating cells and microbial biofilms. This study is the first one to provide a detailed description of the chemical profile of *A. cerefolium* extract dealing with scientific insights into its anti-glioblastoma and antimicrobial activities.

Keywords: *A. cerefolium*; extract; herba; phenolic composition; anti-glioblastoma; antimicrobial; mechanisms of action

1. Introduction

According to the World Health Organization (WHO) grading of central nervous system (CNS) tumors, glioblastoma (GBM) is classified as a malignant grade IV glioma tumor [1]. With poor prognosis and a median survival time of about 15 months GBM is the most aggressive glioma and one of the deadliest forms of brain cancer [2,3]. In most cases by the time of diagnosis GBM is already widely spread. Existing therapeutic strategies for GBM include surgery, radiation and chemotherapy. Combined radiotherapy and chemotherapy are currently used for the cytoreduction of the tumor, but over 90% of patients usually experience rapid tumor recurrence [4–6]. Tumor recurrence, poor prognosis,
and the side effects of radio- and chemotherapy suggest the need to prioritize the further search for novel therapeutic approaches, especially among natural sources.

It has been revealed that microbial infections contribute to cancer promotion through the production of carcinogenic metabolites and alterations in host physiological processes [7,8]. There is cumulative evidence that Candida albicans is able to stimulate the onset and progression of cancers by provoking inflammation and producing carcinogens such as acetaldehyde [9]. On the other side, bloodstream infections, no surgery, more than two hospitalizations, distant metastasis, absence of drainage tubes and radiotherapy are potential prognostic risk factors in cancer patients with S. aureus infection [10]. Therefore, the search for novel natural substances that can be used to prevent or treat cancer and related microbial infections is a high priority.

Anthriscus cerefolium (L.) Hoffm. is an annual aromatic plant, belonging to the chervil plant genus of the family Apiaceae. The genus encompasses twelve species, some of which are considered weeds [11]. For thousands of years, since the times of the ancient Greeks, the delicate young leaves of chervil have been used in spring tonics. Herbalists have used chervil throughout history as a diuretic, expectorant, digestive and skin freshener. It is also thought to relieve the symptoms of eczema, gout, kidney stones and pleurisy. Chervil is a traditional remedy for bad dreams, burns and stomach problems and supposedly, the whole plant alleviates hiccups. It is used as an eyewash for rinsing and refreshing the eyes [11,12]. Previous research regarding chemical composition of Anthriscus has been mainly focused on the flavonoids present in A. sylvestris. Thus quercetin, apigenin and rutin have been identified in A. sylvestris [13]. Furthermore, Dall’Acqua et al. [14] showed that fractions of this plant contain mainly luteolin-7-O-glucoside (cinaroside).

Since herbalists have used chervil throughout history for numerous biological activities, one of the aims of the present study was to investigate the chemical composition of the methanolic extract of herba Anthrisci cerefolii. As traditional uses of the plant include its use against bad dreams, meaning that it probably contains compounds that can pass the blood-brain barrier, we investigated anti-glioblastoma activity of methanolic extract of A. cerefolium. Since the plant was traditionally used for the treatment of pneumonia, suggesting its antimicrobial effect, we tested the activities of extract against microbes Staphylococcus aureus and Candida species.

2. Results and Discussion

2.1. Analysis of Phenolic Acids and Flavonoids

A review of the literature revealed that the chemical compounds derived from Anthriscus cerefolium have not been extensively studied. In the genus Anthriscus, the essential oils [15] and lignans [16] were the most investigated chemical compounds. Here we investigate the presence of phenolic acid and flavonoid derivatives in the methanolic extract of herba Anthrisci cerefolii. Since there is no literature data regarding these chemical compounds in Anthriscus cerefolium, we compared our results with results obtained in some other Anthriscus species, as well as in other plants belonging to the Apiaceae family (Table 1).
Table 1. High resolution mass spectrometry (HRMS) and MS^4 data for phenolic acids and flavonoids identified in *Anthriscus cerefolium* extract.

| Peak No. | Identified Compounds | $t_r$, min | Molecular Formula, [M–H]^− | Calculated Mass, [M–H]^− | Exact Mass, [M–H]^− | Δ ppm | MS^2 Fragments, (% Base Peak) | MS^3 Fragments, (% Base Peak) | MS^4 Fragments, (% Base Peak) | Reference |
|----------|----------------------|-----------|----------------------------|---------------------------|---------------------|------|-----------------------------|-----------------------------|-----------------------------|-----------|
| 1        | Dihydroxybenzoyl hexoside | 3.95      | C_{13}H_{17}O_{5}s        | 315.02716                 | 315.07246           | −0.31| 109(10), 152(22), 153(100), 154(9), 268(10), 278(9), 279(23) | 108(7), 109(100) | NA                          | /         |
| 2        | Caffeoyl-hexoside isomer 1 | 4.61      | C_{15}H_{17}O_{7}s        | 341.08781                 | 341.08750           | 0.31 | 133(4), 179(100), 180(3), 89(19), 129(15), 134(11), 135(100), 143(23), 144(20), 161(13) | 135(100) | 79(21), 107(100), 117(49) | [17]      |
| 3        | Caffeoyl-hexoside isomer 2 | 4.48      | C_{15}H_{17}O_{7}s        | 339.07216                 | 339.07142           | 0.74 | 177(100), 178(3) | 89(4), 105(10), 133(100), 149(6) | 89(105), 105(9), 123(15) | [17]      |
| 4        | Caffeoyl-hexoside isomer 3 | 5.11      | C_{15}H_{17}O_{7}s        | 341.08781                 | 341.08674           | 1.07 | 135(9), 179(100), 180(7), 295(3) | 85(95), 93(56), 111(41), 127(100), 171(32), 173(81) | 83(11), 85(100), 97(10), 99(38), 109(29) | [19]      |
| 5        | Caffeoyl-hexoside isomer 4 | 5.24      | C_{15}H_{17}O_{7}s        | 353.08781                 | 353.08753           | 0.28 | 179(3), 191(100) | 66(100), 91(60) | NA                          | [17]      |
| 6        | Caffeoyl-hexoside isomer 5 | 5.29      | C_{15}H_{17}O_{7}s        | 163.04007                 | 163.03968           | 0.39 | 99(13), 115(18), 116(12), 119(100), 128(16), 131(19), 135(12) | 96(100), 91(60) | NA                          | [17]      |
| 7        | Caffeoyl-hexoside isomer 6 | 5.87      | C_{15}H_{17}O_{7}s        | 337.09289                 | 337.09270           | 0.19 | 163(4), 173(8), 191(100) | 85(95), 93(56), 111(41), 127(100), 171(32), 173(81) | 83(11), 85(100), 97(10), 99(38), 109(29) | [19]      |
| 8        | Caffeoyl-hexoside isomer 7 | 6.32      | C_{15}H_{17}O_{7}s        | 367.10346                 | 367.10372           | 0.74 | 191(100), 192(8), 193(4), 321(4) | 179(4), 191(100) | NA                          | [17]      |
| 9        | Caffeoyl-hexoside isomer 8 | 7.04      | C_{15}H_{17}O_{7}s        | 356.11950                 | 515.11763           | 1.87 | 191(13), 335(9), 55(100), 354(14) | 179(4), 191(100) | NA                          | [21]      |
| 10       | Caffeoyl-hexoside isomer 9 | 7.12      | C_{15}H_{17}O_{7}s        | 601.11989                 | 601.12114           | 1.25 | 395(55), 439(72), 442(12), 515(85), 516(20), 557(100), 558(24) | 233(32), 335(4), 377(9), 395(100) | 173(13), 234(100), 335(8) | [22]      |
| 11       | Caffeoyl-hexoside isomer 10 | 7.34     | C_{15}H_{17}O_{7}s        | 601.11989                 | 601.11856           | 1.33 | 233(10), 395(100), 396(13), 439(9), 515(5), 557(8), 558(10) | 233(12), 234(100), 335(5) | 155(3), 173(100) | [22]      |
| 12       | Caffeoyl-hexoside isomer 11 | 7.47      | C_{15}H_{17}O_{7}s        | 601.11989                 | 601.12119           | 1.30 | 233(10), 395(100), 396(13), 439(9), 515(5), 557(8), 558(10) | 233(30), 335(4), 377(10), 395(100), 515(3) | 173(13), 234(100), 335(7) | [22]      |
| 13       | Caffeoyl-hexoside isomer 12 | 7.79      | C_{15}H_{17}O_{7}s        | 367.10346                 | 367.10270           | 0.75 | 191(100), 192(16), 321(17), 322(9), 323(9), 329(8), 330(13) | 85(99), 93(45), 109(29), 127(100), 171(26), 173(56) | NA                          | [17]      |
| 14       | Caffeoyl-hexoside isomer 13 | 8.17      | C_{15}H_{17}O_{7}s        | 356.11950                 | 352.04422           | 0.66 | 93(100) | 93(100) | NA                          | [18]      |
| 15       | Quercetin              | 6.40      | C_{27}H_{27}O_{6}s        | 609.14611                 | 609.14539           | 0.72 | 225(5), 271(7), 300(37), 301(100), 343(12) | 225(5), 271(7), 300(37), 301(100), 343(12) | 151(77), 179(100), 255(45), 257(13), 271(76), 273(19) | [19]      |
| 16       | 3-O-(6′-rhamnopyl)-glucoside | 6.68     | C_{27}H_{27}O_{6}s        | 447.09329                 | 447.08956           | 3.73 | 285(100), 286(13) | 151(41), 175(100), 199(87), 217(80), 241(97), 243(70) | 119(8), 131(86), 133(20), 147(100), 157(5) | [23]      |
| Peak No. | Identified Compounds | 1<sub>b</sub>, min | Molecular Formula, [M–H]<sup>−</sup> | Calculated Mass, [M–H]<sup>−</sup> | Exact Mass, [M–H]<sup>−</sup> | Δ ppm | MS<sup>2</sup> Fragments, (% Base Peak) | MS<sup>3</sup> Fragments, (% Base Peak) | MS<sup>3</sup> Fragments, (% Base Peak) | Reference |
|---------|----------------------|-------------------|----------------------------------|----------------------------------|---------------------------------|-----|----------------------------------|----------------------------------|----------------------------------|-----------|
| 20      | Kaempferol 3-O-(6′-acetyl)-hexoside<sup>c</sup> | 7.20              | C21H21O12<sup>−</sup> | 489.10385                          | 489.10298                        | 0.87 | 285(100), 286(9), 429(6)      | 151(37), 175(62), 199(68), 217(72), 241(100), 243(59) | 151(37), 175(62), 199(68), 217(72), 241(100), 243(59) | [24]      |
| 21      | Apigenin 7-O-glucoside (Apigetrin)<sup>a,b</sup> | 7.22              | C21H21O11<sup>−</sup> | 431.09837                          | 431.09842                        | 0.05 | 268(11), 269(100), 270(11), 311(3) | 149(30), 181(26), 183(27), 224(26), 225(100), 227(29) | 149(30), 181(26), 183(27), 224(26), 225(100), 227(29) | [17]      |
| 22      | Kaempferol 3-O-rhamnoside<sup>c</sup> | 7.65              | C21H19O10<sup>−</sup> | 431.09837                          | 431.09750                        | 0.87 | 255(6), 284(60), 285(100), 286(7), 327(4) | 256(31), 257(100), 267(45) | 163(75), 185(14), 213(23), 229(10), 239(45) | [17]      |
| 23      | Apigenin 7-O-(6′-acetyl)-hexoside<sup>c</sup> | 8.43              | C21H21O11<sup>−</sup> | 473.10894                          | 473.10899                        | 0.95 | 268(50), 269(100), 270(14), 311(6) | 197(34), 201(24), 225(100) | 169(37), 181(35), 183(39), 196(21), 197(100) | [17]      |
| 24      | Luteolin<sup>a,b</sup> | 8.65              | C15H10O6<sup>−</sup> | 285.04046                          | 285.03874                        | 1.73 | 151(28), 175(72), 197(21), 199(69), 217(53), 241(100), 243(53) | 151(34), 169(100), 179(10), 180(14), 182(5) | 151(34), 169(100), 179(10), 180(14), 182(5) | [25]      |
| 25      | Kaempferol 3-O-(6′-p-coumaroyl)-hexoside<sup>c</sup> | 9.39              | C20H20O13<sup>−</sup> | 593.12307                          | 593.12899                        | 0.18 | 285(100), 286(9), 307(31), 308(4) | 251(100), 213(50), 229(57), 241(42), 243(36), 257(87) | 83(4), 107(100) | [26]      |
| 26      | Apigenin<sup>a,b</sup> | 9.50              | C15H10O7<sup>−</sup> | 269.04555                          | 269.04531                        | 0.23 | 149(45), 151(29), 183(17), 201(27), 225(100), 226(18), 227(18) | 169(13), 180(15), 181(100), 183(27), 196(20), 197(38) | 117(17), 139(25), 152(100), 153(41), 163(7) | [25]      |
| 27      | Kaempferol 3-O-(4′-p-coumaroyl)-rhamnoside<sup>c</sup> | 9.90              | C20H20O12<sup>−</sup> | 577.13515                          | 577.13701                        | 1.86 | 285(100), 286(9), 299(16), 300(17), 315(100), 316(11) | 151(80), 229(38), 241(41), 259(33), 257(100), 267(27) | 163(27), 211(7), 213(16), 229(100), 239(12) | [27]      |
| 28      | Isothamnetin 3-O-(3′-p-coumaroyl)-rhamnoside<sup>c</sup> | 10.10             | C20H20O13<sup>−</sup> | 607.14572                          | 607.14621                        | 3.11 | 284(6), 285(27), 299(16), 300(17), 315(100), 316(11) | 300(100) | / | [27]      |
| 29      | Quercetin 3-O-(2′,6′-di-p-coumaroyl)-rhamnoside<sup>c</sup> | 10.50             | C20H20O16<sup>−</sup> | 755.16176                          | 755.16248                        | 0.72 | 271(3), 285(4), 301(3), 307(6), 409(100), 470(20), 503(5) | 135(11), 161(56), 179(100), 271(22), 307(68) | NA | [28]      |
| 30      | Kaempferol 3-O-(2′,6′-di-p-coumaroyl)-hexoside<sup>c</sup> | 11.08             | C20H19O15<sup>−</sup> | 739.16684                          | 739.16671                        | 0.14 | 285(9), 307(4), 453(100), 454(22), 455(4), 593(4) | 135(10), 161(100), 163(31), 179(65), 289(12), 307(67) | 117(3), 133(100) | [29]      |
| 31      | Kaempferol 3-O-(2′,3′-di-p-coumaroyl)-rhamnoside<sup>c</sup> | 11.54             | C20H20O14<sup>−</sup> | 723.17193                          | 723.16845                        | 3.48 | 285(33), 286(8), 437(100), 438(17), 439(3), 577(3) | 145(100), 163(71), 187(24), 211(14), 273(29), 291(46) | 117(100) | [30]      |
| 32      | Kaempferol 3-O-(2′,4′-di-p-coumaroyl)-hexoside<sup>c</sup> | 11.57             | C20H19O15<sup>−</sup> | 753.18249                          | 753.18286                        | 0.36 | 285(100), 296(9), 315(39), 437(49), 453(12), 467(77), 468(12) | 151(87), 185(47), 213(46), 229(60), 239(42), 257(100) | 189(28), 213(57), 215(13), 229(100), 239(24) | [20]      |

<sup>a</sup> Confirmed by standards; <sup>b</sup> Earlier identified in some *Anthriscus* species; <sup>c</sup> Identified in some other species from Apiaceae family; For the first time identified in Apiaceae family; 1<sub>b</sub>, min—retention time; Δ ppm—mean mass accuracy.
Using liquid chromatography in combination with the hyphenated mass spectrometry technique, we identified 32 compounds in total. To the best of our knowledge, this is the first report in which 32 compounds are identified in this plant. Table 1 lists all identified compounds with retention times and major mass spectrometry characteristics (exacts masses, MS$_2$, MS$_3$, and MS$_4$ fragmentations). Among 32 compounds, 17 of them belong to phenolic acids and related compounds, and the 15 of them are flavonoids aglycones and glycosides. The presence of 10 compounds (caffeic acid—3, aesculin—4, chlorogenic acid—6, p-coumaric acid—7, ferulic acid—8, rutin—18, cynaroside—19, apigetrin—21, luteolin—24, and apigenin—26) was confirmed by comparison with available standards.

Regarding phenolic acid derivatives, in addition to simple acids, hexosides and esters with quinic acid were also found. All compounds, except o-hydroxybenzoic acid (compound 17), belong to the group of hydroxycinnamic acids. Compound 17 found at 8.17 min and 137 m/z showed characteristic MS$_2$ base peak at 93 m/z (generated by the loss of CO$_2$—44 Da) and it has been previously detected in Anthriscus vulgaris Bernh. (Apiaceae) from Algeria [18]. Fragmentation pattern and MS spectra of compound 1 (dihydroxybenzoyl hexoside) is fully consistent with the literature [31]. As far as we know, this compound has not been detected so far in any extract of the plant from the Apiaceae family. Compounds 11 and 16 (eluted at 6.32 and 7.79 min, respectively) were marked as feruloylquinic acid isomers. In this case, we can also speculate about the position of esterification because Shrestha et al. [32] showed the differentiation of all feruloylquinic acid derivatives by MS$_2$ base peaks. Namely, unlike other derivatives, MS$_2$ base peak of 5-O-feruloylquinic acid was found at 191 m/z, so these compounds can be cis and trans isomers. Similarly, compounds 13, 14, and 15 (malonyl-dicaffeoylquinic acid isomers) found at 601 m/z, previously identified in Erigeron breviscapus (Apiaceae) extract, were named according to the available literature data [22].

By analyzing flavonoids, compounds from the subgroup of flavonols and flavones were found. Only two aglycones were detected, luteolin (compound 24) and apigenin (compound 26) and both compounds have already been confirmed as constituents of Anthriscus sylvestris Hoffm. (Apiaceae) [25]. The most common compounds were flavonol glycosides with a p-coumaroyl residue. Almost all such compounds showed specific fragmentation where the mass of deprotected aglycone does not appear as MS$_2$ base peak, but it is formed by loss of mass of aglycone. Thus, for example, compound 31 at 11.54 min and 723 m/z generated MS$_2$ base peak at 437 m/z, which corresponds to the fragment resulting from kaempferol loss [M–H–286]$,^-$, while the secondary MS$_2$ peak was found at 285 m/z (deprotonated kaempferol). Further, MS$_3$ fragmentation gave a base peak at 145 m/z (loss of two p-coumaroyl residues) and secondary peaks at 291 and 163 m/z (Figure 1). Based on all these findings, a given compound is presumed to be kaempferol 3-O-(2''$^,$3''-di-p-coumaroyl)-rhamnoside, which was previously isolated from the flowers of Foeniculum vulgare Mill. and Foeniculum dulce DC. (Apiaceae) [30].

Another compound that, to the best of our knowledge, has not been previously detected in any extract of the plants from the Apiaceae family is isorhamnetin 3-O-(3''-p-coumaroyl)-rhamnoside (28). The exact position of the p-coumaroyl residue can only be assumed, but the only compound corresponding to the exact mass and fragmentation of our compound was found in Persicaria glabra (Willd.) M.Gómez extract [33]. Its proposed structure and fragmentation are shown in Figure 2.
Figure 1. Schematic representation of fragmentation of compound 31 (kaempferol 3-O-(2‴,3‴-di-p-coumaroyl)-rhamnoside).

Figure 2. Proposed fragmentation pattern of compound 28 (isorhamnetin 3-O-(3‴-p-coumaroyl)-rhamnoside).

2.2. Anti-Glioblastoma Activity

Cytotoxic effect of the extract on A172 glioblastoma cell line was tested using crystal violet assay. Obtained result was compared with result achieved on human gingival fibroblast (HGF-1) cells. Results are presented in the Table 2. Extract concentration required for 50% inhibition of growth (IC50) of A172 glioblastoma cell line was 765.21 µg/mL. On the other hand, the extract displays no cytotoxicity on HGF-1 primary cells at concentrations up to 800 µg/mL.

| Cell lines | A. cerefolium IC50 (µg/mL) |
|------------|---------------------------|
| A172       | 765.21 ± 56.7             |
| HGF-1      | >800                      |

Previous investigations showed that A. cerefolium extract had potent cytotoxic effect as determined by brine shrimp lethality assays on Artemia larvae [34]. Aqueous extract of A. sylvestris leaves showed no cytotoxicity towards RAW264.7 macrophages cells, but induced an anti-inflammatory defense response in this cell model [35]. Essential oil of the aerial parts of A. caucalis showed cytotoxic activity on liver hepatocellular carcinoma...
(HepG2) and human breast adenocarcinoma (MCF-7) cells [36]. Our results on *A. cerefolium* are supporting previous data regarding cytotoxicity against cancer cell lines of the species of the genus. However, this is the first report of the cytotoxic effects of *A. cerefolium* methanolic extract against a glioblastoma cell line.

The cell morphology of A172 glioblastoma cells treated with vehicle control or *A. cerefolium* methanolic extract (IC₅₀ concentration) was analyzed by examining the expression of the cytoskeletal protein tubulin by immunofluorescence and laser confocal microscopy (Figure 3). As shown in Figure 3A, A172 control cells treated with vehicle (DMSO) exhibited a characteristic fibroblast-like morphology. The microtubule network is interconnected and appears filamentous (Figure 3A). In contrast, A172 cells treated with *A. cerefolium* methanolic extract (Figure 3B) lost their fibroblast-like morphology became rounded with the fragmented nuclei. They exhibited a diffuse tubulin staining pattern and some of them contained multiple micronuclei (arrowheads in Figure 3B). Detected morphological changes are indicative for mitotic arrest and cell death (apoptosis). Further research is needed in order to dissect mechanism of action of the methanolic extract on A172 morphology.

![Figure 3](image)

*Figure 3.* Morphology of A172 control cells (A) and A172 cells treated with *A. cerefolium* methanolic extract (B). Boxed regions in B are enlarged in (B’, B”) figures. White arrowheads in B mark cells with multiple micronuclei.
The impact of the extract on the proliferation of A172 glioblastoma cells was investigated by analyzing the expression of Ki67 protein, a marker of proliferation. Ki67 is well-known marker for determination of the proliferation of tumor; its expression is used as a prognostic marker for cell proliferation in many tumors and it has been revealed that Ki67 value predicts the response to neoadjuvant chemotherapy [37]. The percentage of proliferating cells (Ki-67 labeling index) can discriminate more aggressive phenotypes of tumors; currently, thus the values of Ki67 labeling index is used both to predict the prognostic stratification of patients and to estimate the responsiveness to the resistance to chemotherapy [38]. By Ki67 immunostaining we detected that proliferation rate of A172 cells treated with the extract was decreased by approximately 30% compared to proliferation rate of control cells (Figure 4). Our study is the first to explore the effect of herba Anthrisci cerefolii on the properties of glioblastoma cells. The obtained results provide a good basis for further research that could explore the possibilities of applying the extract of herba Anthrisci cerefolii in the treatment of glioblastoma.

It has been shown previously that quercetin, catechins and proanthocyanidins inhibit the proliferation of glioblastoma cells and induce their death; being able to cross the blood-brain barrier. Some of them inhibit pro-oncogene signaling pathways and intensify the effect of conventional anti-cancer therapies [39]. Our results indicate that extract of herba Anthrisci cerefolii was rich source of flavonoid molecules.
2.3. Antimicrobial Activities of Herba Anthrisci cerefolii

The antimicrobial activity of the *A. cerefolium* extract was tested by the microdilution method (Table 3). The most sensitive species to the effect of the *A. cerefolium* extract was yeast *C. tropicalis*, while the most resilient species to the microbicidal effect was the *S. aureus* ATCC 11632 bacterium strain. The results obtained for the extract were comparable with results obtained for the positive controls. It is important to highlight that extract was active against methicillin resistant strain of *S. aureus*.

Table 3. Antimicrobial activity of the *A. cerefolium* methanolic extract (mg/mL).

| Bacteria          | *A. cerefolium* | Streptomycin | Ampicillin |
|-------------------|-----------------|--------------|------------|
| *S. aureus* (ATCC 11632) | 2.50           | 0.17         | 0.34       |
| *S. aureus* MRSA  | 1.25           | 0.10         | -          |
| *S. aureus* MRSA  | 2.50           | -            | -          |
| Yeasts            | *A. cerefolium* | *Ketoconazole* | *Bifonazole* |
| *C. albicans* (ATCC 10231) | 1.25           | 0.50         | 0.15       |
| *C. krusei* (clinical isolate) | 1.25           | 1.00         | 0.30       |
| *C. tropicalis* (ATCC 750) | 0.62           | 0.30         | 0.25       |

Previous investigations were focused on essential oils from the genus *Anthriscus* and there are no reports about antimicrobial activity of methanolic extract of *A. cerefolium*. Namely, the essential oil from the aerial parts of *Anthriscus caucalis* M. Bieb showed significant activity against *Bacillus subtilis* and *Escherichia coli* with MIC values of 0.095 mg/mL and 0.105 mg/mL, respectively [36]. The essential oil obtained from the root of *Anthriscus nemorosa* (Bieb.) Sprengel showed significant activity against the following microbes: *Candida albicans*, *Staphylococcus epidermidis*, *Bacillus subtilis* and *Escherichia coli* [40].

The effect of the methanolic extract of *A. cerefolium* on *Staphylococcus aureus* (clinical isolate) biofilm formation was assessed by bacterial biofilm inhibition assay. The inhibition of biofilm formation by *S. aureus* was achieved at sub-MICs of the *A. cerefolium* extract. Bacterial biofilm was inhibited at 1/2 MIC for 69.88%, while at lower MICs inhibition capacity decreased (Table 4A). Regarding the inhibition of pre-formed yeast biofilms, the extract was equally active against all of the tested *Candida* species (Table 4B). Our results strongly point to the antibiofilm potential of the extract. These results are significant since microbial biofilms are more resistant to conventional therapeutics. According to our knowledge, there are no previous reports on antibiofilm potential of *Anthriscus* species.

Table 4. The effects of *A. cerefolium* methanolic extract on: (A) bacterial biofilm formation by *S. aureus* at sub-MICs (%) and (B) on MIC and MFC (mg/mL) in formed fungal biofilm.

### (A) Inhibition of *S. aureus* bacterial biofilm formation

|                     | 1/2 MIC | 1/4 MIC | 1/8 MIC | 1/16 MIC | 1/32 MIC |
|---------------------|---------|---------|---------|----------|----------|
| *A. cerefolium*     | 69.88 ± 6.86 | 67.91 ± 4.99 | 44.25 ± 4.70 | NI       | NI       |
| Streptomycin        | 55.64 ± 2.12 | 35.33 ± 1.47 | 33.22 ± 1.08 | 15.21 ± 1.12 | NI       |

### (B) Inhibitory and fungicidal effects on formed fungal biofilms

| Fungi               | *A. cerefolium* | Fluconazole |
|---------------------|-----------------|-------------|
|                     | MIC             | MFC         | MIC       | MFC       |
| *C. albicans* (ATCC 10231) | 5.00           | 10.00       | 8.00      | 9.00      |
| *C. krusei* (clinical isolate) | 5.00           | 10.00       | 2.00      | 3.00      |
| *C. tropicalis* (ATCC 750) | 5.00           | 10.00       | 3.00      | 6.00      |
Our results indicate that the extract of *A. cerefolium* did not significantly inhibit staphyloxanthin production in *S. aureus* at sub-MICs (data not shown). The extract had no influence on the ergosterol biosynthetic pathway in *C. albicans*, as well. The extract was able to induce leakage of cellular components in *C. albicans* (Figure 5), suggesting that the main mode of antifungal action is in the level of cell membrane permeability. Our results provide the first highlighting of the impact of *A. cerefolium* extract on the yeast cell membrane permeability.

![Graph](image-url)  
**Figure 5.** Leakage of cellular components in *C. albicans* measured at 260 nm and 280 nm.

### 3. Materials and Methods

#### 3.1. Collection and Extraction of Plant Material

*Anthriscus cerefolium* (L.) Hoffm. (Apiaceae) was collected in Belgrade, Serbia, during the flowering period of plant in May 2018. The aerial parts of the plant were lyophilized and reduced to a fine powder. Plant material was successively extracted with methanol according to the procedure described previously [41,42].

#### 3.2. Chemicals and Reagents

Solvents for UHPLC/MS analysis (acetonitrile and formic acid) were of MS grade, obtained from Fisher Scientific (Loughborough, UK). Ultrapure water was generated by deionization (Millipore, Billerica, MA, USA). Standards of phenolic acids and flavonoids (caffeic acid, aesculin, chlorogenic acid, *p*-coumaric acid, ferulic acid, rutin, cynaroside, apigetrin, luteolin, and apigenin) were purchased from Sigma-Aldrich (Steinheim, Germany).

#### 3.3. UHPLC-LTQ Orbitrap MS4 Analysis of Phenolic Acid and Flavonoid Derivatives

Analysis of compounds of interest was carried out by an Accela UHPLC system connected with LTQ Orbitrap mass spectrometer equipped with heated-electrospray ionization (HESI) ionization applied in negative mode (ThermoFisher Scientific, Bremen, Germany). The separation was achieved using a Syncronis C18 column (100 × 2.1 mm, 1.7 µm particle size; ThermoFisher Scientific). The gradient elution program, settings of ion source and the other parameters of the mass detector were the same as previously described [43]. Identification of compounds was done according to their monoisotopic mass (obtained by full scan analysis) and MS3 fragmentation and also confirmed by literature data. Accurate mass of compounds was calculated using ChemDraw software (version 12.0, CambridgeSoft, Cambridge, MA, USA) and for instrument control, data acquisition and data analysis Xcalibur software (version 2.1, Thermo Fisher Scientific, Waltham, MA, USA) was used.

#### 3.4. Investigation of the Antiproliferative Effect of *A. cerefolium* Extract

Crystal violet assay was used to study the cytotoxic effect of *A. cerefolium* extract on A172 glioblastoma cell line and human gingival fibroblasts cells HGF-1 (ATCC® CRL-2014™). Cytotoxic effect of *A. cerefolium* methanolic extract on HGF-1 cells was determined...
as described by Stojkovic et al., [42]. The A172 cells were grown in high-glucose Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 2 mM L-glutamine and 1% penicillin and streptomycin at 37 °C in 10% CO₂. The day before treatment, 4 × 10⁵ cells were seeded per well in a 96-well plate. On the day of treatment, the fresh medium with different concentrations of the extract (250–1000 µg/mL) dissolved in dimethyl sulfoxide (DMSO) was added to the cells and cells were incubated with the extract for 48h. After that period, cells were washed twice with phosphate-buffered saline (PBS), stained with 0.5% crystal violet staining solution for 15 min at room temperature and after removal of crystal violet the cells were washed in a stream of tap water and left to air-dry at room temperature. Later, the dye was dissolved in methanol and the absorbance of dye was measured in a microplate reader Infinite 200 PRO at 590 nm. Control cells were treated with the same percentage of DMSO as treatment with the highest concentration of the extract. The DMSO concentration in the assay did not exceed 0.5%. The experiment was done in triplicate for each concentration of the extract and three independent experiments were performed. The results were expressed as IC50 values in µg/mL.

3.5. Immunocytochemistry

Twenty four hours before treatment 2.5 × 10⁴ A172 cells were seeded per well on coverslips in a 12-well plate. After that, cells were treated with the extract (IC50 concentration) or vehicle DMSO (control cells) for 48 hours. Later, cells were fixed in 4% paraformaldehyde for 15 min at room temperature (RT), washed 3 times for 20 min in 1× PBS, permeabilized 10 min in 0.2% Triton X-100 in PBS and blocked for 1h at RT in 10% normal goat serum/1% bovine serum albumin (BSA) in PBS. Primary antibodies, rabbit anti-Ki67 antibody (diluted 1:250, Abcam, Cambridge, UK) and mouse anti-tubulin antibody (Abcam, diluted 1:100) were diluted in PBS containing 1% BSA/0.1% Triton X-100. After one hour incubation at room temperature with anti-Ki67 antibody, cells were washed 3 times for 15 min with 0.1% Triton X-100 in PBS and incubated with anti-rabbit secondary antibody conjugated with Alexa Fluor 488 (diluted 1:500 in 1% BSA/0.1% Triton X-100 in PBS) (Invitrogen, Life Technologies Corporation, Carlsbad, CA, USA) for 1 h. Afterward, cells were washed 3 times for 15 min with 0.1% Triton X-100 in PBS and stained with 0.1 mg/mL diaminophenylindole (DAPI; Sigma, Saint Louis, MO, USA). Images were taken using a BX51 fluorescent microscope (Olympus, Tokyo, Japan) equipped with appropriate filters and analyzed using the Cytovision software (Applied Imaging Corporation, Santa Clara, CA, USA). Ki67 index was determined as number of Ki67 positive cells (without measurement of fluorescence intensity)/total number of cells. At least 10 independent images per experiment were chosen to score Ki67 positive cells in controls and at least 15 independent images per experiment were chosen to score Ki67 positive cells in treatments with the extract. Images for control and treatment were taken with the same magnification. In other experiments, after overnight incubation at 4°C with mouse anti-tubulin antibody, cells were washed 3 times for 15 min with 0.1% Triton X-100 in PBS and incubated for one hour at RT with biotinylated anti-mouse IgG antibody (diluted 1:500 in 1% BSA/0.1% Triton X100 in PBS, Vector Laboratories, Burlingame, CA, USA). Following washing 3 times for 15 min with 0.1% Triton X-100 in PBS, cells were incubated with DyLight 594 Streptavidin antibody (diluted 1:500 in PBS, Vector Laboratories) for one hour at RT. After washing 3 times for 15 min with 0.1% Triton X-100 in PBS, nuclei were stained with 0.1 mg/mL DAPI. Images were taken using a TCS SP8 confocal microscope (Leica Microsystems, Wetzlar, Germany) applying the Leica LAS AF-TCS SP8 software.

3.6. Antimicrobial Susceptibility Tests

The A. cerefolium methanolic extract at the concentration of 0.1–20 mg mL⁻¹ (in 5% dimethylsulfoxide—DMSO) was tested for antibacterial and antifungal activity through the serial microdilution method as described previously [41,42]. All microorganisms were provided by the Institute for Biological Research “Sinisa Stankovic”—National Institute of the Republic of Serbia, University of Belgrade (Belgrade, Serbia). Staphylococcus aureus
(ATCC 11632), and methicillin-resistant S. aureus (MRSA isolate) were used for antibacterial analysis. For analysis of antifungal activity, the Candida albicans (ATCC 10231), C. tropicalis (ATCC 750), and C. krusei (clinical isolate) were used. Streptomycin and ampicillin were used as positive controls for antibacterial assays, whereas ketoconazole and bifonazole were selected as positive controls for the antifungal assays, while 5% DMSO was used as a negative control. The results were presented as minimum inhibitory concentration (MIC, required for microbial growth inhibition), bactericidal (MBC) and fungicidal concentrations (MFC), expressed in mg/mL.

3.7. Activity against Formation and Inhibition of Microbial Biofilms
The effect of A. cerefolium methanolic extract on Staphylococcus aureus (clinical isolate) biofilm formation was assessed by bacterial biofilm inhibition assay. Crystal violet was used as a staining solution for the biofilm and biofilm formation inhibition was evaluated by spectrophotometric techniques. Results were presented as percentage of biofilm formation inhibition with respect to untreated control [41].

The effect of A. cerefolium methanolic extract on inhibition of formed biofilm of C. albicans, C. krusei and C. tropicalis was determined as previously described [42]. MIC was defined as the minimum concentration of the extract that inhibited further growth of the initial biofilm, and minimum fungicidal concentration (MFC) represents the concentration of the extract that resulted in the level of luminescence presenting no fungal growth.

3.8. Insights into Modes of Antibacterial and Antifungal Actions of Extract
The staphyloxanthin inhibition assay in S. aureus was performed as previously described [41]. SubMIC concentrations of the extract, ranging from 1/2 MIC to 1/32 MIC, were used for the assay. The pigment production was measured at 465 nm using a spectrophotometer. The results were presented as percentage of staphyloxanthin production in treated bacterium with respect to non–treated control bacterium.

A Candida albicans strain was used for ergosterol binding and membrane permeability assays. In order to determine whether the antifungal effect of the extract was achieved via disruption of ergosterol biosynthetic pathway, serial dilutions (same as for microdilution method) of the extract were prepared with addition of ergosterol (25–100 µg/mL). After 24 h of incubation at 37 °C, MFC values were determined as explained for antifungal activity assay. The effect of A. cerefolium extract on cell membrane permeability was analyzed as described previously. The optical density of the filtrate was measured at room temperature (25 °C) at 260 and 280 nm (8453 spectrophotometer, Agilent, Santa Clara, CA 95051 United States) [42].

4. Conclusions
Methanolic extract obtained from herba Anthrisci cerefolii was investigated for the first time regarding its phenolic composition. Phenolic acids and their derivatives, phenolic-related compounds and flavonoids were identified in the extract. The extract decreased the proliferation rate of A172 glioblastoma cells and induced cell morphology changes indicative for mitotic arrest and apoptosis, while at the same time it was not cytotoxic to control HGF-1 cells. Additionally, the extract was active against bacteria and yeast, and the obtained results suggest that antifungal mode of action is associated with disruption of cell membrane permeability. Previously, it was published that some natural products, like green barley extract [44], and lactoferrin [45] exhibited an antiproliferative activity on cancer cells, making our results in agreement with literature pointing to the anticancer effects of natural products. Altogether, the obtained results serve as a good basis for further research of the anti-glioblastoma and antimicrobial mechanisms of actions of herba Anthrisci cerefolii.

Author Contributions: D.S. conceived and designed the research, was enrolled in all the analyses and prepared the draft version of the manuscript; D.D. and M.S. (Marija Schwirtlich) were engaged in the experiments on in vitro cytotoxicity, immunocytochemistry and microscopy, interpreted re-
sults and edited the draft version of the manuscript; N.R. edited the final draft of the manuscript and participated in the conceptualization of the project; M.S. (Milena Stevanović) supervised the experiments on cell line, edited the final draft of the manuscript; M.D.S. supervised the whole project, and edited the final manuscript; U.G. performed chemical analyses, data curation, and methodology, reviewed and edited the final manuscript, supervised the project. All authors have read and agreed to the published version of the manuscript.

**Funding:** This work has been supported by Ministry of Education, Science and Technological Development of Republic of Serbia (451-03-68/2020-14/200007 and 451-03-68/2020-14/200042).

**Data Availability Statement:** All data is presented in the manuscript.

**Conflicts of Interest:** The authors declare no conflict of interest.

**References**

1. Louis, D.N.; Perry, A.; Reifenberger, G.; von Deimling, A.; Figarella-Branger, D.; Cavenee, W.K.; Ohgaki, H.; Wiestler, O.D.; Kleihues, P.; Ellison, D.W. The 2016 World Health Organization Classification of Tumors of the Central Nervous System: A summary. *Drug Discov. Today* 2016, [CrossRef] [PubMed]

2. Luwor, R.B.; Stylli, S.S.; Kaye, A.H. The role of Stat3 in glioblastoma multiforme. *J. Clin. Neurosci.* 2013, 20, 907–911. [CrossRef] [PubMed]

3. Thakkar, J.P.; Dolecek, T.A.; Horbinski, C.; Ostrom, Q.T.; Lightner, D.D.; Barnholtz-Sloan, J.S.; Villano, J.L. Epidemiologic and molecular prognostic review of glioblastoma. *Cancer Epidemiol. Biomark. Prev.* 2014, 23, 1985–1996. [CrossRef] [PubMed]

4. Neagu, M.R.; Reardon, D.A. Rindopepimut vaccine and bevacizumab combination therapy: Improving survival rates in relapsed glioblastoma patients? *Immunotherapy* 2015, 7, 603–606. [CrossRef] [PubMed]

5. Kasl, F.T.; Watts, C. Novel Surgical Approaches to High-Grade Gliomas. *Curr. Treat. Options Neurol.* 2015, 17, 38. [CrossRef] [PubMed]

6. Zanders, E.D.; Svensson, F.; Bailey, D.S. Therapy for glioblastoma: Is it working? *Drug Discov. Today* 2019, 24, 1193–1201. [CrossRef] [PubMed]

7. Eyvazi, S.; Vostakolaei, M.A.; Dilmaghani, A.; Borumandi, O.; Nezhad, M.S.; Hahroba, H.; Tarhriz, V. The oncogenic roles of bacterial infections in development of cancer. *Microb. Pathog.* 2020, 141, 104019. [CrossRef]

8. Chang, A.H.; Parsonnet, J. Role of bacteria in oncogenesis. *Clin. Microbiol. Rev.* 2010, 23, 837–857. [CrossRef] [PubMed]

9. Ramirez-Garcia, A.; Remeterea, A.; Aguirre-Urizar, J.M.; Moragues, M.D.; Antoran, A.; Pellon, A.; Habad-Diaz-De-Cerio, A.; Hernando, F.L. *Candida albicans* and cancer: Can this yeast induce cancer development or progression? *Crit. Rev. Microbiol.* 2016, 42, 181–193. [CrossRef]

10. Bai, C.; Li, D.; Zhang, Q.; Zheng, S.; Li, Z.; Wang, H.; Li, L.; Zhang, W. Prognostic analysis of cancer patients with *Staphylococcus aureus* infection: Five-year experience at a comprehensive cancer center. *Int. J. Clin. Exp. Med.* 2018, 11, 8640–8645.

11. Vyas, A.; Shukla, S.S.; Pandey, R.; Jain, V.; Joshi, V.; Gidwani, B. Chervil: A multifunctional miraculous nutritional herb. *Asian J. Plant Sci.* 2012, 11, 163–171. [CrossRef]

12. Farooqi, A.A.; Srivivasappula, K.N. Chervil. In *Handbook of Herbs and Spices*, 2nd ed.; Elsevier Inc.: Amsterdam, The Netherlands, 2012; Volume 2, pp. 268–274. ISBN 9780857095688.

13. Milovanovic, M.; Picuric-Jovanovic, K.; Vucelic-Radovic, B.; Vrbski, Z. Antioxidant effects of flavonoids of *Anthriscus sylvestris* in lard. *JAOCS J. Am. Oil Chem. Soc.* 2012; Volume 2, pp. 268–274. ISBN 9780857095688.

14. Dall’Acqua, S.; Giorgetti, M.; Cervellati, R.; Innocenti, G. Deoxypodophyllotoxin content and antioxidant activity of aerial parts of *Anthriscus sylvestris* Hoffm. *Z. Für Nat. C* 2006, 61, 658–662. [CrossRef] [PubMed]

15. Chizzola, R. Composition of the essential oils from *Peucedanum cervaria* and *P. alisticum* growing wild in the Urban Area of Vienna (Austria). *Nat. Prod. Commun.* 2012, 7, 1515–1518. [CrossRef]

16. Hendrawati, O.; Woerdenbag, H.J.; Michiels, P.J.A.; Aantjes, H.G.; Van Dam, A.; Kayser, O. Identification of lignans and related compounds in *Anthriscus sylvestris* by LC-ESI-MS/MS and LC-SPE-NMR. *Phytochemistry* 2011, 72, 2172–2179. [CrossRef]

17. Zengin, G.; Sinan, K.I.; Ak, G.; Mahomoodally, M.F.; Paksoy, M.Y.; Picot-Allain, C.; Glamocilja, J.; Sokovic, M.; Jekö, J.; Cziak, Y.; et al. Chemical profile, antioxidant, antimicrobial, enzyme inhibitory, and cytotoxicity of seven Apiaceae species from Turkey: A comparative study. *Ind. Crops Prod.* 2020, 153, 112572. [CrossRef]

18. Sekhara, I.; Benaisa, O.; Amrani, A.; Giangiacomo, B.; Benabderrahmane, W.; Chaouch, M.A.; Zama, D.; Benayache, S.; Benayache, F. Antioxidant activity and chemical constituents of *Anthriscus vulgaris* Bernh. (Apiaceae) from Algeria. *Acta Sci. Nat.* 2020, 7, 59–70. [CrossRef]

19. Karakaya, S.; Koca, M.; Sytar, O.; Dursunoglu, B.; Ozbek, H.; Duman, H.; Guvenalp, Z.; Kilic, C.S. Antioxidant and anti-cholinesterase potential of *Ferulafo cassisia* with farther bio-guided isolation of active coumarin constituents. *S. Afr. J. Bot.* 2019, 121, 536–542. [CrossRef]

20. de la Luz Cádiz-Gurrea, M.; Fernández-Arroyo, S.; Joven, J.; Segura-Carretero, A. Comprehensive characterization by UHPLC-ESI-Q-TOF-MS from an *Eryngium bourgatii* extract and their antioxidant and anti-inflammatory activities. *Food Res. Int.* 2013, 50, 197–204. [CrossRef]
21. Bouratoua, A.; Khalfallah, A.; Bensouici, C.; Kabouche, Z.; Alabud Magid, A.; Harakat, D.; Voutquenne-Nazabadioko, L.; Kabouche, A. Chemical composition and antioxidant activity of aerial parts of Ferula longipes Coss. ex Bonnier and Maury. Nat. Prod. Res. 2018, 32, 1873–1880. [CrossRef]

22. Zhang, Y.; Shi, P.; Qu, H.; Cheng, Y. Characterization of phenolic compounds in Erigeron breviscapus by liquid chromatography coupled to electrospray ionization mass spectrometry. Rapid Commun. Mass Spectrom. 2007, 21, 2971–2984. [CrossRef] [PubMed]

23. Žemlička, L.; Fodran, P.; Lukeš, V.; Vagánek, A.; Slováková, M.; Staško, A.; Dubaj, T.; Liptaj, T.; Karabin, M.; Birošová, L.; et al. Physicochemical and biological properties of luteolin-7-O-β-d-glucoside (cynaroside) isolated from Anthriscus sylvestris (L.) Hoffm. Mon. Fur Chem. 2014, 1307–1318. [CrossRef]

24. Spinola, V.; Castilho, P.C. Phytochemical Profile, Chemotaxonomic Studies, and In Vitro Antioxidant Activities of Two Endemism from Madeira Archipelago: Melanoselium decipiens and Monizia edulis (Apiaceae). Chem. Biodivers. 2016, 13, 1290–1306. [CrossRef] [PubMed]

25. Kuríthara, T.; Kikuchi, M. Studies on the constituents of Anthriscus sylvestris Hoffm. II. On the components of the flowers and leaves. Yakugaku Zasshi 1979, 99, 602–606. [CrossRef] [PubMed]

26. Li, D.; Yue, D.; Liu, D.; Liu, X.; Song, S. Chemical constituents from Bupleurum chinense and their chemotaxonomic significance. Biochem. Syst. Ecol. 2019, 86, 103929. [CrossRef]

27. Nazemiyeh, H.; Delazar, A.; Movahedin, N.; Jodari, M.; Imani, Y.; Ghahramani, M.-A.; Nahar, L.; Sarker, S.D. Free radical scavengers from the aerial parts of Grammosciadium platycarpum Boiss. & Hausskn. (Apiaceae) and GC-MS analysis of the essential oils from its fruits. Rev. Bras. Farmacogn. 2019, 19, 914–918. [CrossRef]

28. Abou El-Kassem, L.; Hawas, U.; Awad, H.; Taie, H. Flavonoids from the aerial parts of Ononis spinosa L. Is an antifungal agent with no cytotoxicity to primary human cells. Yakugaku Zasshi 2017, 137, 7138–7151. [CrossRef]

29. Zhang, Z.; Li, S.; Ownby, S.; Wang, P.; Yuan, W.; Zhang, W.; Scott Beasley, R. Phenolic compounds and rare polyhydroxylated terpenoid saponins from Eryngium yuccifolium. Phytochemistry 2008, 69, 2070–2080. [CrossRef] [PubMed]

30. Soliman, F.M.; Shehata, A.H.; Khaleel, A.E.; Ezzat, S.M. An acylated kaempferol glycoside from flowers of Foeniculum vulgare and F. dulce. Molecules 2002, 7, 245–251. [CrossRef]

31. Zengin, G.; Cvetanović, A.; Gašić, U.; Dragićević, M.; Stupar, A.; Uysal, A.; Šenkardes, I.; Sinan, K.I.; Picot-Allain, M.C.N.; Ak, G.; et al. UHPLC-LTQ Orbitrap MS analysis and biological properties of Origanum vulgare subsp. viridulum obtained by different extraction methods. Ind. Crops Prod. 2020, 154, 112747. [CrossRef]

32. Shrestha, A.; Hakeem Said, I.; Grimbs, A.; Thielen, N.; Lansing, L.; Schepker, H.; Kuhnert, N. Determination of hydroxycinnamic acids present in Rhododendron species. Phytochemistry 2017, 144, 216–225. [CrossRef] [PubMed]

33. Manivannan, R.; Shopna, R. Isolation of Quercetin and Isorhamnetin Derivatives and Evaluation of Anti-microbial and Anti-inflammatory Activities of Persicaria glabra. Nat. Prod. Sci. 2015, 21, 170–175.

34. Jovanova, B.; Kulevanova, S.; Kadíjkova Panovska, T. Determination of the total phenolic content, antioxidant activity and cytotoxicity of selected aromatic herbs. Agric. Conspec. Sci. 2019, 84, 51–58.

35. Lee, S.A.; Moon, S.-M.; Han, S.H.; Hwang, E.J.; Hong, J.H.; Park, B.-R.; Choi, M.S.; Ahn, H.; Kim, J.-S.; Kim, H.-J.; et al. In Vivo and in Vitro Anti-Inflammatory Effects of Aqueous Extract of Anthriscus sylvestris Leaves. J. Med. Food 2018, 21, 585–595. [CrossRef] [PubMed]

36. Lai, P.; Rao, H.; Gao, Y. Chemical composition, cytotoxic, antinflammatory and antioxidant activities of essential oil from Anthriscus caucalis M. Bieb grown in China. Rec. Nat. Prod. 2018, 12, 290–294. [CrossRef]

37. Pascale, M.; Aversa, C.; Barbazza, R.; Marongiu, B.; Siracusano, S.; Stoffel, F.; Sulfarico, S.; Roggero, E.; Bonin, S.; Stanta, G. The proliferation marker Ki67, but not neuroendocrine expression, is an independent factor in the prediction of prognosis of primary prostate cancer patients. Radiol. Oncol. 2016, 50, 313–320. [CrossRef] [PubMed]

38. Armocida, D.; Pesce, A.; Di Giammarco, F.; Frati, A.; Salvati, M.; Santoro, A. Histological, molecular, clinical and outcomes characteristics of Multiple Lesion Glioblastoma. A retrospective monocentric study and review of literature. Neurocirugía 2020. [CrossRef] [PubMed]

39. Vidak, M.; Rozman, D.; Komel, R. Effects of flavonoids from food and dietary supplements on glial and glioblastoma multiforme cells. Molecules 2015, 20, 19406–19432. [CrossRef]

40. Pavlović, M.; Petrović, S.; Milenković, M.; Couladis, M.; Tzakou, O.; Niketić, M. Chemical composition and antimicrobial activity of Anthriscus nemorosus root essential oil. Nat. Prod. Commun. 2011, 6, 271–273. [CrossRef]

41. Stojković, D.; Drakulić, D.; Gašić, U.; Zengin, G.; Stevanović, M.; Ražićević, N.; Soković, M. Ononis spinosa L., an edible and medicinal plant: UHPLC-LTQ-Orbitrap/MS chemical profiling and biological activities of the herbal extract. Food Funct. 2020, 11, 7138–7151. [CrossRef]

42. Stojković, D.; Dias, M.I.; Drakulić, D.; Barros, L.; Stevanović, M.; Ferreira, I.C.F.R.; Soković, M.D. Methanolic extract of the herb Ononis spinosa L. Is an antifungal agent with no cytotoxicity to primary human cells. Pharmaceuticals 2020, 13, 78. [CrossRef] [PubMed]

43. Banjana, T.; Dragićević, M.; Šiler, B.; Gašić, U.; Bohane, B.; Nestorović Živković, J.; Trifunović, S.; Mišić, D. Chemodiversity of two closely related tetraploid Centaureum species and their hexaploid hybrid: Metabolomic search for high-resolution taxonomic classifiers. Phytochemistry 2017, 140, 27–44. [CrossRef] [PubMed]
44. Robles-Escajeda, E.; Lerma, D.; Nyakeriga, A.M.; Ross, J.A.; Kirken, R.A.; Aguilera, R.J.; Varela-Ramirez, A. Searching in Mother Nature for Anti-Cancer Activity: Anti-Proliferative and Pro-Apoptotic Effect Elicited by Green Barley on Leukemia/Lymphoma Cells. *PLoS ONE* **2013**, *8*, e73508. [CrossRef] [PubMed]

45. Nakamura-Bencomo, S.; Gutierrez, D.A.; Robles-Escajeda, E.; Iglesias-Figueroa, B.; Siqueiros-Cendón, T.S.; Espinoza-Sánchez, E.A.; Árvalo-Gallegos, S.; Aguilera, R.J.; Rascón-Cruz, Q.; Varela-Ramirez, A. Recombinant human lactoferrin carrying humanized glycosylation exhibits antileukemia selective cytotoxicity, microfilament disruption, cell cycle arrest, and apoptosis activities. *Investig. New Drugs* **2020**. [CrossRef]