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The Anti-Acne Potential and Chemical Composition of Two Cultivated Cotoneaster Species

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Abstract: In light of current knowledge on the role of reactive oxygen species and other oxidants in skin diseases, it is clear that oxidative stress facilitates inflammation and is an important factor involved in skin diseases, i.e., acne. Taking into consideration the fact that some Cotoneaster plants are valuable curatives in traditional Asian medicine, we assumed that thus far untested species C. hsingshangensis and C. hissaricus may be a source of substances used in skin diseases. The aim of this study was to evaluate the antioxidant, anti-inflammatory, antimicrobial, and cytotoxic activities of their various extracts. LC-MS analysis revealed the presence of 47 compounds (flavonoids, phenolic acids, coumarins, sphingolipids, carbohydrates), while GC-MS procedure allowed for the identification of 42 constituents (sugar derivatives, phytosterols, fatty acids, and their esters). The diethyl ether fraction of C. hsingshangensis (CHs-2) exhibited great ability to scavenge free radicals and good capacity to inhibit cyclooxygenase-1, cyclooxygenase-2, lipoxygenase, and cyclooxygenase-2, lipoygenase, and hyaluronidase. Moreover, it had the most promising power against microaerobic Gram-positive strains, and importantly, it was non-toxic toward normal skin fibroblasts. Taking into account the value of the calculated therapeutic index (>10), it is worth noting that CHs-2 can be subjected to in vivo study and constitutes a promising anti-acne agent.

Keywords: Cotoneaster; Rosaceae; antioxidant; anti-inflammatory; anti-acne; skin diseases; antimicrobial

1. Introduction

The results obtained by Sarici et al. [1] concerning the evaluation of parameters associated with oxidative stress, such as nitric oxide (NO), xanthine oxidase (XO), malondialdehyde (MDA), superoxide dismutase (SOD), and catalase (CAT), in the venous blood of patients using spectrophotometrical methods indicate that oxidative damage of tissues is a significant etiological factor of acne. In view of the above, the role of oxidative stress in the etiology of acne seems to be unquestionable.

The cutaneous propionibacteria (P. acnes, P. avidum, P. granulosum, P. propionicum, and P. lymphophilum) are involved in the maintenance of healthy skin; however, they can also reveal adverse activity as opportunistic pathogens [2]. The three predominant genera, including Propionibacteria, Staphylococcic, and Corynebacteria, form the microbial community established on the skin [3].

According to [4], the classification of Propionibacterium is extremely complicated. The major types of Propionibacterium (I, II, III clades) should be classified as P. acnes subsp. acnes, P. acnes subsp. defendens, and P. acnes subsp. elongatum, respectively. The latest research justifies the necessity to divide the genus Propionibacterium into four genera. Additionally,
the cutaneous Propionibacterium, participating in seborrheic skin disorders, should be reclassified into the new genus *Cutibacterium*. Nevertheless, the previous classification is still valid and practiced among the medical microbiology community [4]. In our article, the new nomenclature (*C. acnes*) was used.

Although the association of acne lesion aggravation with the presence of *C. acnes* is well established, the mechanism of involvement of *C. acnes* in the formation of acne is not fully understood [5]. Nevertheless, it is well known that *C. acnes* participates in the production of pro-inflammatory cytokines, such as interleukin (IL)-8, IL-1β, IL-12, and tumor necrosis factor (TNF)-α, by stimulating keratinocytes, as well as phagocytic cells, which consequently leads to severe inflammation [5]. Furthermore, according to [6], enhancement of the mRNA expression levels of certain cytokines, such as (IFN)-γ, TNF-β, IL-8, IL-4, IL-10, IL-1α, IL-1β, and IL-17A, within acne lesions can be observed.

The presence of *C. acnes* on the skin leads to the formation of superoxide anions, which form peroxidants after combination with nitric oxide. As a consequence, this phenomenon causes keratinocyte damage [7].

Although synthetic antioxidant compounds are considered very effective, they also cause substantial side effects. Herbal sources possess the ability to participate in various phases of the oxidation mechanism. According to Soleymani and co-authors [8], phenolic compounds show significant anti-acne activity due to their ability to decrease oxidative stress. Many mechanisms of action are arranged in this biological activity and involve activation of TLR-2/4, p38/JNK/MAPK pathways, adjustment of MAPK signaling pathways, and regulation of Nrf2/Keap1-mediated antioxidant pathways. Additionally, upregulation of SOD, GP x, GSH, CAT, and heme oxygenase HO-1 was noted. Furthermore, polyphenolic compositions demonstrate the capability to downregulate endothelial ROS, NOX-4, and MDA (malondialdehyde) levels, which consequently leads to reduced amounts of H₂O₂ and MDA and to the inhibition of the expression of MAPKs, such as P38, ERK, and JNK. Pathways of particular significance, such as the ROS/MAPK/NF-κB pathway and PI3K/Akt/NF-κB pathway, are being suppressed. The decrease in lipid peroxidation linked with the downregulation of cytochrome P450 (CYP) expression and upregulation of STAT-1 2E1 expression seem to be notably relevant.

Medicinal plants have been traditionally used in the treatment of several human diseases, and their therapeutic properties have been assigned to different chemical compounds isolated from them. Of considerable importance, compounds with antioxidant properties may be found at great concentrations in plants and can be responsible for their preventive effects in various diseases caused by oxidative stress. Therefore, the antioxidant activities of plant extracts have prospective applications in healthcare [9].

A diversity of occurrence among the *Cotoneaster* genus is worth underlining and encompasses river valleys and banks, woods, thickets, rocky, and calcareous sites, as well as mountain areas approximately 800–4100 m above sea level. The individual species are indigenous, particularly to the Palearctic region (temperate Asia, Europe, and North Africa). Nevertheless, they are cultivated not only throughout Europe but also in various parts of the world due to their significant aesthetic qualities: white, red, or pink flowers and red, brownish red, orange, or black fruits. The pivotal area of occurrence, with 60% of species, can be observed in mountain regions of China [10]. *Cotoneaster hsingshangensis* is indigenous to China, and *C. hissaricus* is vernacular throughout Asia. These two species, similar to many others of *Cotoneaster*, are cultivated in Poland as ornamental plants in parks, gardens, and other urban places.

It has been reported that many *Cotoneaster* species have valuable biological properties and have been used in traditional medicine in many countries [11]. Swati and co-authors [12] reported that fruits of *Cotoneaster microphyllus* are applied on skin against irritation. Furthermore, leaves of *C. microphyllus* are used for dermatitis. The medicinal uses of *Cotoneaster* species in skin disorders have been reported in Pakistan, Turkey, India, Lebanon, and Iran as therapeutic agents in the treatment of cuts and wounds [13]. In addi-
tion, an anti-itching effect of Cotoneaster has been observed [14,15]. According to Akbar [16], Cotoneaster removes hyperpigmentation of the skin.

Taking into consideration the fact that some representatives of Cotoneaster are valuable curatives applied in various skin diseases in traditional Asian medicine, we assumed that thus far untested species—C. hsingshangensis and C. hissaricus—may be a source of active substances used in skin diseases. In view of the above, the present study reports primarily on a thorough examination of the composition (especially phenolic components) and biological activities with an emphasis on dermatological diseases, such as acne. An exploration of Cotoneaster species has been undertaken to provide an explanation for their capability to ameliorate skin conditions.

2. Materials and Methods

2.1. Chemicals and Reagents

Ascorbic acid, 2,2-diphenyl-1-picrylhydrazyl radical (DPPH•), 2,2′-azino-bis-(3-ethylbenzothiazole-6-sulfonic acid) (ABTS••), (−)-epigallocatechin gallate (EGCG), nordihydroguaiaretic acid (NDGA), indomethacin, hyaluronidase from bovine tests, hyaluronic acid sodium salt from rooster comb, Folin–Ciocalteu reagent, ethylene-diaminetetraacetic acid, disodium dihydrate (Na2EDTA*2H2O), Tricine (≥99%; titration) were obtained from Sigma-Aldrich (Steinheim, Germany). Phosphate-buffered saline (PBS) was purchased from Gibco (Carlsbad, CA, USA). Reference substances were supplied by ChromaDex (Irvine, CA, USA), while acetonitrile, formic acid, and water were supplied for LC analysis by Merck (Darmstadt, Germany). All other chemicals were of analytical grade and were obtained from the Polish Chemical Reagent Company (POCH, Gliwice, Poland).

2.2. Leaves of Cotoneaster hsingshangensis Fryer et B.Hylmö (Figure 1), as well as C. hissaricus Pojark. (Figure 2) were collected in the Maria Curie-Skłodowska University (UMCS) Botanical Garden in Lublin (Poland), at an altitude of 181.2 m a.s.l. (coordinates 52°14′34.4″ N 17°05′32.4″ E and 46°57′51.7″ N 142°45′22.1″ E, respectively) in September 2020. Taxonomical identification was confirmed by Dr. A. Cwener, an employee of the Botanical Garden who specializes in Cotoneaster. C. hsingshangensis was introduced to cultivation in the Botanical Garden in Lublin in 1966. This species was derived from the Institute of Dendrology of the Polish Academy of Sciences, Kôrnîk Arboretum. C. hissaricus was introduced to the Botanical Garden in Lublin in 2003 from Sakhalin Botanical Garden. Voucher specimens were deposited in the Department of Pharmaceutical Botany (CHs-0921 and CHi-0921).

Figure 1. Cont.
2.3. Preparation of the Extracts

The plant materials were dried in the shade at 24 °C (±0.5 °C) to achieve a constant weight [17]. Extracts were prepared using a mixture of methanol–acetone–water (3:1:1, v/v/v; 3 × 100 mL), and then sonicated at a controlled temperature (40 ± 2 °C) for 30 min [13]. The combined extracts were filtered, concentrated under reduced pressure, and after freezing, lyophilized in a vacuum concentrator (Free Zone 1 apparatus; Labconco, Kansas City, KS, USA) to obtain dried residues. The obtained extracts were dissolved in hot water, filtered after 24 h, and subjected to liquid-liquid extraction with diethyl ether, ethyl acetate, and n-butanol, successively. The obtained fractions of diethyl ether, ethyl acetate,
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and n-butanol, as well as the water residue, were evaporated in vacuo and lyophilized using a vacuum concentrator.

2.4. Total Flavonoid, Phenolic, and Phenolic Acids Content

Total flavonoid (TFC) and total phenolic content (TPC) were established using colorimetric assays as described previously [18]. The absorbance was measured at 430 and 680 nm, respectively, using a Pro 200F Elisa Reader (Tecan Group Ltd., Männedorf, Switzerland). TPC was estimated from the calibration curve (R² = 0.9845), using gallic acid as a standard (concentration ranged 0.002–0.1 mg/mL). The results were expressed as mg of gallic acid equivalent (GAE) per 1 g of dry extract (DE). TFC was estimated from the calibrated curve (R² = 0.995), using quercetin (0.004–0.11 mg/mL) as a standard. The results were expressed as mg of quercetin equivalent (QE) per 1 g of DE. Total phenolic acid (TPAC) content was assessed using Arnow’s reagent as described in the Polish Pharmacopoeia IX (an official translation of PhEur 7.0) [17]. The absorbance was measured at 490 nm. TPAC was estimated from the calibration curve (R² = 0.9999), using caffeic acid as a standard in a concentration of 3.36–23.52 µg/mL. The results were expressed as mg of caffeic acid equivalent (CAE) per 1 g of DE.

2.5. LC-MS Analysis

The chromatographic measurements were performed using the LC/MS system from Thermo Scientific (Q-EXACTIVE and ULTIMATE 3000, San Jose, CA, USA) equipped with an ESI source. The ESI was operated in negative polarity modes under the following conditions: spray voltage—3.5 kV; sheath gas—40 arb. units; auxiliary gas—10 arb. units; sweep gas—10 arb. units; and capillary temperature—320 °C. Nitrogen (>99.98%) was employed as sheath, auxiliary, and sweep gas. The scan cycle used a full-scan event at a resolution of 60,000. A Gemini C18 column (4.6 × 100 mm, 3 µm) (Phenomenex, Torrance, CA, USA) was employed for chromatographic separation, which was performed using gradient elution. Mobile phase A was 25 mM formic acid in water; mobile phase B was 25 mM formic acid in acetonitrile. The gradient program started at 5% B, increasing to 95% for 60 min, followed by isocratic elution (95% B) for 10 min. The total run time was 70 min at the mobile phase flow rate 0.4 mL/min. The column temperature was 25 °C. In the course of each run, MS spectra in the range of 100–700 m/z were collected continuously. Additionally, the MS2 functions were used to carry out a detailed qualitative analysis. The collision energy for each examined compound was 25%.

The amounts of the identified compounds were carried out based on the calibration curves obtained for the standard. In the case of quantitative analysis of compounds that do not have standards, calibration curves for substances of similar structure were used. All the results are presented as the mean of three independent measurements (n = 3).

2.6. GC-MS Analysis

The qualification of the sample extract was performed using a GC-MS/MS system (GCMS-TQ8040; Shimadzu, Kyoto, Japan) equipped with a ZB5-MSi fused-silica capillary column (30 m × 0.25 mm i.d., 0.25 µm film thickness; Phenomenex, Torrance, CA, USA). Grade 5.0 helium was used as the carrier gas. Column flow was 1 mL/min. The injection of a 1 µL sample was performed using an AOC-20i + s type autosampler (Shimadzu, Kyoto, Japan). The injector was working at a temperature of 310 °C. The following temperature program was applied: the oven temperature was held at 60 °C for 2 min and was subsequently increased linearly at a rate of 6 °C/min to 310 °C, where it was held for 15 min. The mass spectrometer was operated in EI mode at 70 eV; the ion source temperature was 225 °C. The mass spectra were measured in the range 40–450 amu. The amounts of the individual analytes were estimated by the peak normalization method.
2.7. Antioxidant Activity

All antioxidant and enzyme inhibitory assays were done in 96-well plates (Nunc, Nunc, Roskilde, Denmark) using Infinite Pro 200F Elisa Reader (Tecan Group Ltd., Männedorf, Switzerland). The experiments were performed in triplicate.

2.7.1. DPPH• Assay

The 2,2-diphenyl-1-picryl-hydrazyl (DPPH•) free radical scavenging activity of Cotoneaster extracts and the positive control—ascorbic acid (AA)—was studied using the method described previously [18], but with some modifications. After 30 min of incubation at 28 °C, the decrease in DPPH• absorbance, caused by the tested extracts, was measured at 517 nm. The results were expressed as values of IC50.

2.7.2. ABTS•+ Assay

The ABTS•+ decolorization assay was the second method applied for the assessment of antioxidant activity [18]. The absorbance was measured at 734 nm. Trolox was used as a positive control. The results were expressed as values of IC50.

2.7.3. Metal Chelating Activity (CHEL)

The metal chelating activity was established using the method described by Guo et al. [19], modified in our previous study [18,20]. The absorbance was measured at 562 nm. As a positive control, Na2EDTA*2H2O was used. Results were expressed as the IC50 values of the Cotoneaster extracts based on concentration-inhibition curves.

2.8. Enzyme Inhibitory Activity

2.8.1. Cyclooxygenase-1 (COX-1) and Cyclooxygenase-2 (COX-2) Inhibitory Activity

The extracts of Cotoneaster species were examined for cyclooxygenase-1 (COX-1) and cyclooxygenase-2 (COX-2) inhibitory activity using a COX (ovine/human) Inhibitor Screening Assay Kit (Cayman Chemical, MI, USA) according to the protocol of the manufacturer. The extracts were tested at different concentrations. Indomethacin was used as a positive control.

2.8.2. Lipoxygenase Inhibitory Activity

Anti-lipoxygenase activity of Cotoneaster extracts was determined using the Lipoxygenase Inhibitor Screening Assay Kit (Cayman Chemical, MI, USA) according to the protocol of the manufacturer. The extracts were tested at different concentrations. The effective concentration (µg/mL) in which lipoxygenase activity is inhibited by 50% (IC50) was estimated graphically. Nordihydroguaiaretic acid (NDGA) was used as a positive control.

2.8.3. Hyaluronidase Inhibitory Activity

Anti-hyaluronidase activity was established using the method described by Liyanarrachchi et al. [21]. After 20 min incubation at 37 °C, the absorbance was measured at 585 nm. The extracts were tested at different concentrations. Epigallocatechin gallate was used as a positive control.

2.9. Bacterial Strains

The antibacterial power of Cotoneaster extracts was determined using bacterial strains causing skin diseases (including seborrhea and acne). We used microaerobic Gram-positive bacteria: Cutibacterium granulosum PCM 2462, C. acnes PCM 2334, C. acnes PCM 2400, (possessed from the Polish Collection of Microorganisms PCM Institute of Immunology and Experimental Therapy Polish Academy of Sciences, Poland, as Propionibacterium, now—Cutibacterium), C. acnes ATCC 11827; aerobic Gram-positive: Staphylococcus epidermidis ATCC 12228 and S. aureus ATCC 25923; and aerobic Gram-negative strains Pseudomonas aeruginosa ATCC 27853 and Escherichia coli ATCC 25992. Each bacterial strain was pre-incubated overnight at 37 °C on agar plates. Mueller–Hinton (BioMaxima S.A., Lublin,
Poland) agar or broth (MH-agar, MH-broth) for aerobic strains and Brain–Heart Infusion (Oxoid Ltd, Altrincham, England) agar or broth (BHI-agar, BHI-broth) for microaerobic bacteria were used. The bacteria were suspended in 5 mL of sterile saline water, and the absorbance of this inoculum was adjusted to 108 CFU/mL (0.5 McFarland scale).

2.10. Disc Diffusion Assay

This disc diffusion assay can evaluate the antibacterial potency of tested extracts and was made according to proven methods [22,23]. Briefly, solid medium in Petri dishes was smeared with inoculum of 0.5 McFarland. The plant extracts were dissolved in DMSO (10 mg/mL) then loaded over sterile filter paper discs (8 mm in diameter) to obtain a final concentration of 0.1 mg per disc. Inoculated plates with samples were incubated at 37 °C for 24 h (aerobic stains) or 48 h (microaerobic bacteria).

The zones of growth inhibition around plant samples were measured [mm] and recognized as antibacterial potency. The larger the zone of growth inhibition, the greater the antibiotic activity.

2.11. Minimum Inhibitory Concentration (MIC) and Minimum Bactericidal Concentration (MBC) Determination

The MIC is the lowest concentration of a test compound that inhibits microbial proliferation. The test examined the MIC of the Cotoneaster extracts against Staphylococcus aureus ATCC 25923, S. epidermidis ATCC 12228, C. acnes ATCC 11827, C. acnes PCM 2334, C. acnes PCM 2400, and C. granulosum PCM 2462, whose optical density was 0.5 McFarland scale. Double microdilution in the 96-well plate assay was used according to the CLSI method with some modification, as described in our paper [23,24]. Such an amount of the extract solution to the wells was added, that using the principle of double microdilution to obtain final concentrations in the range of 3.9–1000 µg/mL. Then, each well was inoculated (2 µL of inoculum density of 0.5 McFarland). A background control (broth with extract), a negative control (the broth alone), and a positive control (broth and inoculum) were also prepared. The plates were incubated under conditions proper to bacterial growth (aerobic bacteria at 37 °C, for 24 h; anaerobic bacteria at 37 °C, for 48 h). Finally, microbial growth was determined using a BioTek Synergy H4 (BioTek, Winooski, VT, USA) plate reader at a wavelength of 600 nm.

MBC is the lowest concentration of the agent with bactericidal properties. To determine this, 96-well plates obtained in the previous experiment were used to determine the MIC of the tested extracts. Ten µL of solutions taken from the wells in which no bacterial growth was observed were applied to the new Petri plates. Then, the plates were incubated under optimal conditions for the growth of the given bacteria. After the incubation was completed, a visual assessment of the agar surface of the plates was made. The MBC was considered to be the one in which there was no visible bacterial growth on the solid medium.

To visualize the obtained results, it is advantageous to present the MBC/MIC ratio. Thus, MBC/MIC values ≤4 demonstrate that the agent is bactericidal. MBC/MIC values >4 indicate the bacteriostatic nature of the tested agents [25].

Microbiological tests were performed in three separate experiments (n = 3).

2.12. Cytotoxic Activity

This experiment was carried out according to the protocol described by us previously [26]. The cytotoxic activity of selected substances was evaluated toward the BJ cell line (normal human fibroblasts, ATCC CRL-2522TM). Briefly, the BJ cells were seeded in 96-well plates, and then after 24 h incubation at 37 °C in suitable conditions (5% CO2, 95% air), the culture medium was replaced with two-fold serial dilutions of investigated substances (1.95–1000 µg/mL). After 24 h incubation, the cell viability was assessed using the MTT assay. The obtained data were presented as mean values ± standard deviations (SD). These results were subjected to four-parameter nonlinear regression analyses (GraphPad Prism 5,
version 5.04, GraphPad Software, San Diego, CA, USA) in order to determine values of half-maximum cytotoxic concentration (CC₅₀).

2.13. Statistical Analysis

The results were expressed as mean values ± standard deviation (SD) of three independent experiments. The data from cell culture experiments were subjected to statistical analysis using unpaired Student’s t-test or One-Way ANOVA test, followed by a Tukey’s multiple comparison test, and differences were considered significant when p < 0.05 (GraphPad Prism 5, version 5.04, GraphPad Software, San Diego, CA, USA), whereas Principal Component Analysis was carried out in R version 3.6.3 (64-bit, Windows 10), using built-in “prcomp” function.

3. Results

3.1. Phytochemical Analysis

Total phenolic content (TPC) for Cotoneaster extracts and fractions was determined using Folin-Ciocalteu reagent, and the results were estimated as gallic acid equivalents (GAE) per g of dry extract (DE) (Table 1). Our results showed that the leaves of Cotoneaster (CHi) have the highest phenolic content (296.13 ± 1.52 mg GAE/g DE) than C. hsingshangensis (CHs) (193.84 ± 1.14 mg GAE/g DE). The results obtained in our study were better compared to data presented for extracts derived from the leaves of other Cotoneaster species. For instance, Kicel et al. [27] demonstrated that TPC for the 70% aqueous methanolic extracts of leaves of different Cotoneaster species cultivated in Poland varied from 51.7 (C. tomentosus) to 154.3 mg GAE/g (C. bullatus) of dry weight of the plant material. The same authors [28] also found lower amounts of phenolic compounds in 70% aqueous methanolic extracts of the fruits (from 26 to 43.5 mg GAE/g PM) of different Cotoneaster species.

### Table 1. The total content of phenolic (TPC), flavonoid (TFC), and phenolic acids (TPAC) in the C. hissaricus and C. hsingshangensis leaf extracts.

| Sample | Extraction Yield (% DE) | Total Phenolic Content [mg GAE/g DE] | Total Phenolic Acids [mg CAE/g DE] | Total Flavonoid Content [mg QE/g DE] |
|--------|-------------------------|--------------------------------------|-----------------------------------|-------------------------------------|
| CHi    | 15.30                   | 193.84 ± 1.14 ± e,f,g,h,i             | 33.80 ± 1.03 ± e,f,g,h,i          | 25.38 ± 0.35 ± e,f,g,h,i           |
| CHi-1  | 3.00                    | 83.87 ± 0.23 ± a,e,f,g,h,i            | 10.31 ± 0.11 ± a,e,f,g,h,i        | 0.12 ± 0.05 ± d,f,g,h,i            |
| CHi-2  | 0.89                    | 348.05 ± 2.81 ± a,b,c,d,e,f,g,h,i     | 40.57 ± 0.19 ± a,b,c,d,e,f,g,h,i  | 72.12 ± 0.35 ± a,b,c,d,e,f,g,h,i   |
| CHi-3  | 17.29                   | 219.00 ± 0.49 ± a,b,c,d,e,f,g,h,i     | 37.83 ± 0.15 ± a,b,c,d,e,f,g,h,i  | 58.71 ± 0.49 ± a,b,c,d,e,f,g,h,i   |
| CHi-4  | 6.86                    | 463.16 ± 3.94 ± a,b,c,d,e,f,g,h,i     | 56.12 ± 0.27 ± a,b,c,d,e,f,g,h,i  | 134.89 ± 0.51 ± a,b,c,d,e,f,g,h,i  |
| CHs    | 27.22                   | 296.13 ± 1.52 ± e                     | 61.27 ± 0.93 ± e                  | 47.72 ± 0.37 ± e                   |
| CHs-1  | 6.73                    | 143.30 ± 1.09 ± e                     | 19.65 ± 0.30 ± e                  | 0.23 ± 0.02 ± f                    |
| CHs-2  | 2.07                    | 486.04 ± 3.17 ± e,f                   | 83.94 ± 0.25 ± f                  | 97.66 ± 0.18 ± e,f                 |
| CHs-3  | 14.04                   | 381.83 ± 2.53 ± e                     | 70.17 ± 0.13 ± e                  | 146.58 ± 1.10 ± e                 |
| CHs-4  | 5.76                    | 599.77 ± 3.76 ± e                     | 91.95 ± 0.48 ± e                  | 252.27 ± 0.24 ± e                 |

DE—dry extract; GAE—Gallic Acid Equivalent; CAE—Caffeic Acid Equivalent; QE—Quercetin Equivalent; CHi—methanol–acetone–water (3:1:1, v/v) extract of C. hissaricus; CHs—methanol–acetone–water (3:1:1, v/v) extract of C. hsingshangensis; CHi-1—water fraction of C. hissaricus; CHi-2—diethyl ether fraction of C. hissaricus; CHi-3—butanol fraction of C. hissaricus; CHi-4—ethyl acetate fraction of C. hissaricus. Values were presented as mean ± standard deviation (n = 9). Statistical analysis: a—significantly different results compared to CHi; b—significantly different results compared to CHi-1; c—significantly different results compared to CHi-2; d—significantly different results compared to CHi-3; e—significantly different results compared to CHs; f—significantly different results compared to CHs-1; g—significantly different results compared to CHs-2; h—significantly different results compared to CHs-3; i—significantly different results compared to CHs-4; One-Way ANOVA test, followed by a Tukey’s multiple comparison test, p < 0.05.

In our study, we also examined the content of phenolics in fractions obtained after fractionating crude extracts between solvents of different polarity. The results showed that the TPC level was in the range of 83.87 ± 0.23 (water fraction of C. hissaricus) to 559.77 ± 3.76 mg GAE/g DE (ethyl acetate fraction of C. hsingshangensis). Kicel and co-
authors [29] obtained higher values with the highest TPC level for ethyl acetate fractions (470.9–650.8 mg GAE/g dw) and diethyl-ether fractions (453.1–546.9 mg GAE/g dw) of defatted methanol extracts.

Thus, the results obtained in our study indicated that the leaves of C. hissaricus and C. hsingshangensis are a rich source of phenolic compounds.

The total flavonoid content of the leaves of C. hissaricus and C. hsingshangensis was estimated by a previously described colorimetric method [27]. The data were expressed as quercetin equivalents (QE) per g of dry extract (DE). The results presented in Table 1 show that the content of total flavonoids in both species is at an average level. The comparable content was noted for the leaves of C. hissaricus (CHi) and C. hsingshangensis (CHs) (25.38 ± 2.35 and 47.72 ± 0.37 mg QE/g DE, respectively). The results obtained in our study were higher than those found by Mahmutović-Dizdarević et al. [11]. In their study, quantitative estimation revealed that methanolic extracts from leaves of C. tomentosus possessed 18.17 ± 0.30 mg QE/g dw. flavonoid content, followed by C. integerrimus—16.42 ± 0.35 mg QE/g dw. and C. horizontalis—10.55 ± 0.51 mg QE/g dw. Moreover, they found that fruits of these three species contain a lower amount of flavonoids—2.76–9.38 mg QE/g dw. A higher amount of flavonoids was found in the methanolic extract of leaves of C. wilsonii Nakai (36.46 ± 1.89 mg QE/g dw), while in the stems and fruits, the content of flavonoids was lower (6.09 ± 0.71 and 0.23 ± 0.20 mg QE/g dw, respectively) [30].

Kicel et al. [29] also obtained lower results for defatted methanol extracts of the three Cotoneaster species cultivated in Poland, and the values were from 14.70 ± 0.05 to 67.71 ± 0.80 mg/g dw. These authors obtained slightly higher results for the crude fractions compared to the results in our study. The highest level of flavonoids was determined for the ethyl acetate fraction of C. integerrimus leaves (403.56 ± 7.48 mg/g dw). In our study, the lowest amount of total flavonoids was found in the methanolic extract of leaves of C. hissaricus (0.12 ± 0.05 and 0.23 ± 0.02 mg QE/g dw, respectively) and the highest was in the ethyl acetate fraction of C. hsingshangensis (252.27 ± 0.24 mg QE/g DE).

The total phenolic acid content (TPAC) in Cotoneaster extracts is presented in Table 1. As for the content of polyphenols and flavonoids, a higher content of phenolic acids was noted for crude extract (61.27 ± 0.93 mg CAE/g DE) and fractions (19.65 ± 0.30–91.95 ± 0.48 mg CAE/g DE) of C. hsingshangensis.

The use of plants in the cure of different diseases depends on their phytochemical composition. Our research of the presence of active compounds in the crude methanol–acetone–water (3:1:1, v/v/v) extracts and different fractions (diethyl ether, ethyl acetate, butanol, and aqueous residual) indicated that methanol–acetone–water extracted the greatest range of constituents from the leaves of both Cotoneaster species. Therefore, in our study, we used only the crude extracts for the LC-MS and GC–MS analyses.

Thus, in the next step of our study, the chemical composition of the extracts obtained from Cotoneaster extracts was investigated using the LC-MS method. Table 2 shows 47 identified compounds, including their molecular formula, theoretical and experimental molecular mass, both errors in ppm and mDa, and the fragments. In our study, flavonoids, phenolic acids, coumarins, cyanogenic glycosides, sphingolipids, and carbohydrates were identified using LC-MS analysis. The chromatograms with marked main peaks are displayed in Figures 3 and 4. The results of the quantitative analysis are presented in Table 3. The amounts of the identified compounds were carried out based on the calibration curves obtained for the standard. In the case of quantitative analysis of compounds that do not have standards, calibration curves for substances of similar structure were used.
Table 2. High-resolution mass spectrometry (HR-MS) of [M − H]⁻ ion and MS2 data.

| Peak No. | Name of Compound | [M − H]⁻ | MS2 | Theoretical Mass [M − H]⁻ (Da) | Experimental Mass [M − H]⁻ (Da) | Δ ppm | Δ mDa | Elemental Composition |
|----------|------------------|---------|-----|-------------------------------|---------------------------------|-------|-------|----------------------|
| 1        | Mannitol         | 181     | 59,71,73,85,89,101,113,119,163 | 181.07122                      | 181.07118                      | 0.22  | ~0.04 | C₄H₁₃O₅               |
| 4        | Quercitin-3-O-(2',3'-O-xylosyl)galactoside | 595 | 271,301b,435 | 595.12992 | 595.12999 | 0.12 | 0.07 | C₆H₁₀O₁₅              |
| 5        | Quercitin-3-O-gentiobioside | 625 | 271,301b,463 | 625.14048 | 625.14038 | 0.16 | ~0.10 | C₆H₁₀O₁₅              |
| 6        | Vitexin-2'-O-arabinoside | 563 | 283,432b | 563.14009 | 563.14011 | 0.14 | ~0.08 | C₆H₁₀O₁₅              |
| 7        | Apigenin-6,8-C-diceloside | 593 | 325,386b,387 | 593.15065 | 593.15077 | 0.20 | 0.12 | C₆H₁₀O₁₅              |
| 8        | Vitexin-2'-O-rhamnoside | 577 | 283,432b | 577.15574 | 577.15571 | 0.05 | ~0.03 | C₆H₁₀O₁₅              |
| 9        | Quercitin-3-O-glucoside (Isoquercitrin) | 463 | 271,300b,301 | 463.08766 | 463.08775 | 0.19 | 0.09 | C₆H₁₀O₁₅              |
| 10       | Quercitin-3-O-galactoside (Hyperoside) | 463 | 271,300b,301 | 463.80766 | 463.80763 | 0.06 | ~0.03 | C₆H₁₀O₁₅              |
| 11       | Kaempferol-3-O-glucoside (Astragalin) | 447 | 294b,300 | 447.09274 | 447.09299 | 0.34 | 0.15 | C₆H₁₀O₁₅              |
| 12       | Quercitin-3-O-rhamnoside (Quercitin) | 447 | 294b,300 | 447.09274 | 447.09271 | 0.07 | ~0.03 | C₆H₁₀O₁₅              |
| 13       | 7-Methylkaempferol-4'-O-glucoside | 461 | 298b,315 | 461.10839 | 461.10849 | 0.22 | 0.10 | C₆H₁₀O₁₅              |
| 14       | 3',4'-dihydroxy-6-methoxyflavone-7-O-rhamnoside | 429 | 255,283b,400,401 | 429.11856 | 429.11853 | 0.07 | ~0.03 | C₆H₁₀O₁₅              |
| 15       | Apigenin-8-glucoside (Vitexin) | 431 | 283,311b,312,341 | 429.11856 | 429.11842 | 0.33 | ~0.14 | C₆H₁₀O₁₅              |
| 16       | Apigenin-7-O-glucoside | 447 | 269,270,431 | 431.09783 | 431.09797 | 0.32 | 0.14 | C₆H₁₀O₁₅              |
| 17       | Biochanin A-7-O-glucoside (Sissootrin) | 445 | 132,211,223,224,239b,240,267 | 445.11348 | 445.11355 | 0.16 | 0.07 | C₆H₁₀O₁₅              |
| 18       | 5,7,2',5'-tetrahydroxyflavanone-7-O-glucoside | 449 | 266b,302 | 449.10839 | 449.10825 | 0.31 | ~0.14 | C₆H₁₀O₁₅              |
| 19       | 5-Methylgenistein-4'-O-glucoside | 445 | 255,283b,417,430 | 445.11348 | 445.11339 | 0.20 | ~0.09 | C₆H₁₀O₁₅              |
| 20       | Gentisic acid 2-O-glucoside (Orcularin) | 315 | 153b,271 | 315.07161 | 315.07149 | 0.38 | ~0.12 | C₆H₁₀O₁₅              |
| 21       | Caffeoylmalic acid | 295 | 133,135,176 | 295.0454 | 295.04533 | 0.24 | ~0.07 | C₆H₁₀O₁₅              |
| 22       | Prunasin | 294 | 161b | 294.09777 | 294.09771 | 0.20 | ~0.06 | C₆H₁₀O₁₅              |
| 23       | Amygdalin | 456 | 89,119,143,179,221,263,323b | 456.15059 | 456.15047 | 0.26 | ~0.12 | C₆H₁₀O₁₅              |
| 24       | 7,8-dimethoxy-6-hydroxycumarin | 221 | 191,206b,207 | 221.045 | 221.04506 | 0.27 | 0.06 | C₆H₁₀O₁₅              |
| 25       | Cotonoa A | 291 | 121,165,247b | 291.15964 | 291.15975 | 0.38 | 0.11 | C₆H₁₀O₁₅              |
| 26       | Horizontoate A | 263 | 148,164,219b | 263.12834 | 263.12852 | 0.68 | 0.18 | C₆H₁₀O₁₅              |
| 27       | 3,3',4'-tri-O-methyllellagic acid | 343 | 255,284,299b,329 | 343.0454 | 343.04554 | 0.41 | 0.14 | C₆H₁₀O₁₅              |
| 28       | Scopoletin | 191 | 104,105,120,148b,176 | 191.03444 | 191.03439 | 0.26 | ~0.05 | C₆H₁₀O₁₅              |
| 29       | Arbutin | 271 | 71,101,108b,109,113,161 | 271.08178 | 271.08175 | 0.11 | C₆H₁₀O₁₅              |
| 30       | 5-Methylgenistein | 283 | 239b,255,269 | 283.06065 | 283.06053 | 0.42 | ~0.12 | C₆H₁₀O₁₅              |
| 31       | Erinodictyol | 287 | 151,162b | 287.05557 | 287.05551 | 0.21 | ~0.06 | C₆H₁₀O₁₅              |
| 32       | 5,7,2',5'-tetrahydroxyflavanone | 287 | 151,162b,259 | 287.05557 | 287.05564 | 0.24 | 0.07 | C₆H₁₀O₁₅              |
| 33       | Naringenin | 271 | 119,151b,187 | 271.06065 | 271.06061 | 0.15 | ~0.04 | C₆H₁₀O₁₅              |
| 34       | Horizontoate C | 482 | 210,272b,288 | 482.42093 | 482.42097 | 0.08 | 0.04 | C₆H₁₀O₁₅              |

b—base peak.
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Figure 3. TIC chromatogram of C. hissaricus extract.

Figure 4. TIC chromatogram of C. hsingshangensis extract.

Table 3. Content of active compounds in the leaves of C. hissaricus (CHi) and C. hsingshangensis (CHs).

| No | Compound | Calibration Standard | Amounts [µg/g DE] |
|----|----------|----------------------|------------------|
|    |          |                      | CHi               | CHs               |
| 1  | mannitol | glucose              | 6834 ± 249        | 3104 ± 123 *     |
| 2  | ascorbic acid | ascorbic acid | 298 ± 10          | 1726 ± 66 *      |
| 3  | quercetin 3-O-rutinoside (rutin) | rutin | 18,028 ± 650       | 3823 ± 131 *     |
| 4  | quercetin 3-O-(2'-O-xylosyl)galactoside | rutin | 7318 ± 289         | 2667 ± 99 *      |
| 5  | quercetin 3-O-gentiobioside | rutin | 2759 ± 109         | 2310 ± 87        |
| 6  | apigenin 6,8-C-dicellobioside | rutin | 5625 ± 233         | 3249 ± 121 *     |
| 7  | vitexin 2''-O-arabinobioside | rutin | 5926 ± 225         | 1273 ± 50 *      |
| 8  | quercetin 3-O-glucoside (isoquercitrin) | rutin | 3923 ± 154         | 1268 ± 49 *      |
| 9  | quercetin 3-O-galactoside (hyperoside) | rutin | 10,079 ± 353       | 8926 ± 327 *     |
| 10 | kaempferol 3-O-glucoside (astragalin) | rutin | 9119 ± 331         | 6184 ± 237 *     |
| 11 | quercetin 3-O-rhamnoside (quercitrin) | rutin | 2480 ± 92          | 4430 ± 168 *     |
| 12 | kaempferol 4'-O-glucoside | rutin | 2158 ± 80          | 6726 ± 251 *     |
| 13 | 3',4'-dihydroxy-6-methoxyflavone | rutin | 3939 ± 137.9       | 2079 ± 79 *      |
| 14 | 7-O-rhamnoside | rutin | 1538 ± 50          | 443 ± 16 *       |
| 15 | apigenin 8-C-glucoside (vitexin) | rutin | 2454 ± 94          | 1930 ± 77 *      |
| 16 | apigenin 7-O-glucoside | rutin | 186 ± 57           | 3734 ± 169 *     |
| 17 | biochanin A 7-O-glucoside (sissotrin) | rutin | 724 ± 26           | 709 ± 23         |
| 18 | 5,7,2',5'-tetrahydroxyflavanone | rutin | 8067 ± 290         | 2469 ± 93 *      |
| 19 | 7-O-glucoside | rutin | 1205 ± 43          | 943 ± 34 *       |
| 20 | 5-methylgenistein 4'-O-glucoside | quercetin | 3838 ± 154        | 1818 ± 79 *      |
| 21 | p-hydroxybenzoic acid | p-hydroxybenzoic acid | 897 ± 30          | 823 ± 27 *       |
| 22 | benzoic acid | benzoic acid | 339 ± 12          | 441 ± 16         |
| 23 | gentisic acid | gentisic acid | 258 ± 10          | <LOQ             |
| 24 | protocatechic acid | protocatechic acid | 509 ± 20          | 69 ± 3 *         |
| 25 | syringic acid | syringic acid | 753 ± 31          | 1130 ± 39 *      |
| 26 | vanillyl acid | vanillyl acid | <LOQ              | <LOQ             |
| 27 | caffeoylmalic acid | caffeic acid | 2828 ± 109        | 1459 ± 51 *      |
Among flavonoids, the most abundant in both species were quercetin derivatives. In the leaves of *C. hissaricus* rutin (18,028 ± 650 µg/g DE), isooroside (10,079 ± 353 µg/g DE), hyperoside (9119 ± 331 µg/g DE), 5,7,2′,5′-tetrahydroxyflavanone 7-O-glucoside (8067 ± 290 µg/g DE), and quercetin 3-O-(2′′-O-xylosyl)galactoside (7318 ± 289 µg/g DE) were found in the largest amount. In the leaves of *C. hsingshangensis*, isooroside (8926 ± 327 µg/g DE), quercetin (6726 ± 251 µg/g DE), and hyperoside (6184 ± 237 µg/g DE) were observed in the greatest amount. Isoquercetin, rutin, hyperoside, and quercetin were previously identified as dominant in the other *Cotoneaster* species [27,28,31–37]. Vitexin 2′-O-arabinoside, vitexin-2′-O-rhamnoside, and 5-methylgenistein, which were found in large quantities in both studied species, were previously noticed only in leaves of *C. thymaefolia* [32], aqueous alcoholic extracts of leafy twigs of *C. obricularis* [38], and chloroform extracts of *C. simonsii* leafy twigs [39], respectively. Subsequent compounds of rare occurrence in the *Cotoneaster* genus are 5,7,2′,5′-tetrahydroxyflavanone and its 7-O-glucoside, which were identified as dominant in *C. thymaefolia* leaves [32]. It is worth noting that rare in nature flavonoids, such as biochanin A 7-O-glucoside (sissotrin) and 5-methylgenistein-4′-O-glucoside, were previously observed in the methanolic extract of leaves of *C. mongolica* [37], methanol extract of *C. serotina* flowers [40], and methanol extract of flowers and fruits of *C. pannosa* [40]. 5-Methylgenistein-4′-O-glucoside was previously observed only in the chloroform extract of leafy twigs of *C. simonsii* [39]. Quercetin 3-O-gentiobioside (quercetin 3-O-β-D-glucopyranosyl(1-6)glucopyranoside) is noteworthy regarding the fact that it was one of the first compounds isolated from the *Cotoneaster* genus and is infrequent, being only reported in *C. oligantha* stems [41]. Furthermore, kaempferol 3-O-glucoside (astragalin) is of rare occurrence in the *Cotoneaster* genus and reported only in the methanol extract of *C. mongolica* leaves by Odontuya and co-authors [37].

The second largest group of active compounds were phenolic acids, and among them, chlorogenic acid, gentisic acid 2-O-glucoside, and caffeoylmalic acid were the most abundant in both species. Apart from these typical phenolic acids, cotonooate A and horizontoate A were observed in the leaves of *C. hissaricus* and *C. hsingshangensis*, but only in *C. hissaricus* was cotonooate A identified in quantifiable amounts (1564 ± 55 µg/g DE). In previous research, cotonooate A and horizontoate A were isolated only from the leafy

### Table 3. Cont.

| No | Compound                          | Calibration Standard | Amounts [µg/g DE] |
|----|-----------------------------------|----------------------|------------------|
|    |                                   |                      | CHi  | CHs  |
|    |                                   |                       | 28   | 29   |
| 28 | chlorogenic acid                  | chlorogenic acid      | 37,932 ± 1330    | 60,043 ± 2231 |
|    |                                   | glucose              | 3360 ± 127      | 1226 ± 61  |
|    |                                   | p-coumaric acid      | 329 ± 13        | 539 ± 19 * |
|    |                                   | glucose              | 1803 ± 63       | 555 ± 19 * |
|    |                                   | caffeic acid         | 6118 ± 223      | 2220 ± 81 *|
|    |                                   | cinnamic acid        | 924 ± 31        | 2429 ± 89 *|
|    |                                   | ferulic acid         | 644 ± 24        | 1567 ± 65 *|
|    |                                   | salicylic acid       | 657 ± 27        | 823 ± 30 * |
|    | 7,8-dimethoxy-6-hydroxycoumarin   | umbelliferone         | 1209 ± 43       | <LOQ      |
|    | cotonooate A                      | benzoic acid         | 1564 ± 55       | <LOQ      |
|    | horizontoate A                    | benzoic acid         | <LOQ            | <LOQ      |
|    | 3,3′,4′-tri-O-methyllellagic acid | quercetin            | 2053 ± 81       | 1169 ± 45 *|
|    | scopoletin                        | umbelliferone        | 12,219 ± 440    | 10,481 ± 371 *|
|    | arbutin                           | glucose              | 1126 ± 41       | 1084 ± 46 *|
|    | 5-methylgenistein                 | quercetin            | 1109 ± 39       | 1423 ± 59 *|
|    | quercetin                         | quercetin            | 229 ± 7         | 115 ± 4 *  |
|    | horizontoate C                    | oleic acid           | 5516 ± 219      | 2939 ± 119 *|
|    | eriodictyol                       | quercetin            | 59 ± 3          | <LOQ      |
|    | 5,7,2′,5′-tetrahydroxyflavanone   | quercetin            | 89 ± 5          | <LOQ      |
|    | naringenin                        | 2627 ± 97           | 2121 ± 88 *     |

LOQ—limit of quantification; DE—dry extract. Asterisk (*) denotes significantly different data (p < 0.05, unpaired *t*-test) compared to CHi.
twigs of the *C. racemiflora* chloroform soluble fraction of the methanolic extract [14] and methanolic extract of *C. horizontalis* [15], respectively.

Chlorogenic acid seems to be a ubiquitous ingredient among the Cotoneaster genus. What most attracts attention is that both studied extracts of leaves of *C. hissaricus* and *C. hsingshangensis* respectively. Chlorogenic acid seems to be a ubiquitous ingredient among the Cotoneaster genus. What most attracts attention is that both studied extracts of leaves of *C. hissaricus* and *C. hsingshangensis* constitute a significant percentage of this acid (37,932 ± 1330 and 60,043 ± 2231 µg/g DE, respectively). According to [42], this compound participates in diminishing *P. acnes*-induced matrix metalloproteinase-9 levels, obstructing nuclear factor-κB (NF-κB) activation, inactivating mitogen-activated protein kinases (MAPK), and decreasing the migration of neutrophils and interleukin (IL)-1β+ populations in vivo. Therefore, it is increasingly being recognized as possibly efficacious in the management of dermatological conditions, such as acne.

Among the other polyphenolic compounds, some coumarins were identified in the leaves of *C. hissaricus* and *C. hsingshangensis*, and the most abundant in both species was scopoletin (12,219 ± 440 and 10,481 ± 371 µg/g DE, respectively). This compound was isolated earlier from the leafy twigs of *C. racemiflora* from methanolic extract [43] and the ethyl acetate-soluble fraction of methanolic extract [44,45].

In both studied species, great amounts of mannitol were also observed (*C. hissaricus*—6834 ± 249 µg/g DE and 3104 ± 123 µg/g DE—*C. hsingshangensis*). This compound is the predominant polysaccharide of manna, which is produced by the young shoots of *C. discolor, C. nummularius, C. tricolor*, and *C. nummularioides* [46]. From cyanogenic glycosides, prunasin, and amygdalin were found. These glycosides were also previously identified in the fruits and leaves of *C. congesta, C. praecox*, and *C. integerrimus* [47] and in the ethanol extract of *C. horizontalis* leafy twigs [48].

Between phenolic compounds occurring in *C. hsingshangensis* and *C. hissaricus*, caffeic acid [44,45], ferulic acid [49,50], chlorogenic acid [51], cinnamic acid [52], quercetin [53], and vitexin [54] are substances with proven anti-acne properties on skin.

The appropriate GC-MS procedure allowed for the identification of 42 and 41 compounds in the leaves of *C. hissaricus* (Table 4, Figure 5) and *C. hsingshangensis* (Table 5, Figure 6), respectively. Among the identified compounds, four main groups of analytes can be distinguished—sugar derivatives, phytosterols, fatty acids, and their esters.

### Table 4. Composition of the extracts of leaves of *C. hissaricus* (% of total fraction; mass%, GC).

| No. | tr    | Area % | Compound                                      |
|-----|-------|--------|-----------------------------------------------|
| 1   | 9.776 | 39557931 2.52 | Benzofuran, 2,3-dihydro-          |
| 2   | 10.908 | 12341078 0.79 | Isosorbide                                 |
| 3   | 12.913 | 3452024 0.22 | Furan, 3-(4-methyl-5-trans-phenyl-1,3-oxazolidin-2-yl)-   |
| 4   | 13.952 | 192528086 12.29 | 1,3,2,5-Dimethylene-1-rhamnitol        |
| 5   | 14.242 | 135536063 8.65 | D-Glycero-D-galacto-heptose             |
| 6   | 15.48  | 7430877 0.47 | Megastigmatriene                                            |
| 7   | 15.988 | 32418863 2.07 | 3-Hydroxy-β-damascone                              |
| 8   | 16.329 | 229108049 14.62 | 2-Deoxy-D-galactose                           |
| 9   | 17.009 | 21401978 1.37 | 2-Hydroxyhexadecyl butanoate               |
| 10  | 17.229 | 21671810 1.38 | (E)-2,6-Dimethoxy-4-(prop-1-en-1-yl)phenol        |
| 11  | 17.448 | 5862283 0.37 | Myristic acid methyl ester                       |
| 12  | 17.709 | 6588889 0.42 | 9-(3,3-Dimethylxiran-2-yl)-2,7-dimethylmna-2,6-dien-1-ol |
| 13  | 18.137 | 7027919 0.45 | 6-Hydroxy-4,4,7a-trimethyl-5,6,7a-tetrahydrobenzofuran-2(4H)-one |
| 14  | 18.951 | 19840196 1.27 | Neophytadiene                                         |
| 15  | 19.034 | 6060805 0.39 | 2-Pentadecanone, 6,10,14-trimethyl-     |
| 16  | 19.222 | 20107664 0.65 | [1,1′-Bicyclopropyl]-2-octanoic acid, 2′-hexyl-, methyl ester |
| 17  | 19.496 | 10732207 0.69 | 3,7,11,15-Tetramethyl-2-hexadecen-1-ol          |
| 18  | 20.028 | 15776294 10.07 | Hexadecanoic acid, methyl ester             |
| 19  | 20.525 | 95409334 6.09 | Palmitic acid                                 |
| 20  | 21.372 | 11475712 0.73 | Cyclohexanebutanoic acid                     |
| 21  | 21.913 | 3242083 0.21 | Methyl trans-4-(2-nonylcylopentyl)butanoate   |
| 22  | 22.029 | 10523560 0.67 | cis-11,14-Eicosadienoic acid, methyl ester    |
| 23  | 22.104 | 33503994 2.14 | Linolenic acid methyl ester                 |
| 24  | 22.17  | 5141879 0.33 | 9-Octadeconic acid (Z), methyl ester         |
| 25  | 22.251 | 247573718 15.79 | Phytol, acetate                             |
| 26  | 22.387 | 53091621 3.39 | Methyl stearate                              |
Table 4. Cont.

| No. | Area | % Area | Compound                          |
|-----|------|--------|-----------------------------------|
| 27  | 23.854 | 32049624 | 0.05 Benzyl beta-D-glucoside      |
| 28  | 24.561 | 3681746  | 0.24 Methyl 18-methyl nonadecanoate |
| 29  | 24.852 | 4060228  | 0.26 4,8,12,16-Tetramethylheptadecan-4-olide |
| 30  | 25.205 | 2015357  | 0.13 3,7,11,15-Tetramethylhexadec-2-en-1-yl acetate |
| 31  | 26.197 | 5666720  | 0.36 1-Heptacosanol               |
| 32  | 26.403 | 17982618 | 1.15 1-Docosanol, methyl ether    |
| 33  | 27.053 | 3938497  | 0.25 Phytol palmitate            |
| 34  | 28.102 | 4304975  | 0.27 9-Hexacosene                 |
| 35  | 28.403 | 2176024  | 0.14 Methyl 18-methyl nonadecanoate |
| 36  | 31.436 | 1813814  | 0.12 Cholesta-4,6-dien-3-ol (3.beta)- |
| 37  | 31.551 | 8162207  | 0.52 Octatriacontyl trifluoroacetate |
| 38  | 31.851 | 40546560 | 2.59 dl-.alpha.-Tocopherol        |
| 39  | 33.331 | 54433136 | 3.47 beta-Sitosterol              |
| 40  | 33.87  | 1158500  | 0.08 Urs-12-en-28-al              |
| 41  | 34.019 | 3680922  | 0.23 Acetyl betulinaldehyde       |

Table 5. Composition of the extracts of leaves of *C. hsingshangensis* (% of total fraction; mass%, GC).

| No. | tr   | Area | % Area | Compound                          |
|-----|------|------|--------|-----------------------------------|
| 1   | 9.655 | 1862168 | 0.19  Benzofuran, 2,3-dihydro-    |
| 2   | 10.990 | 2840045 | 0.28  Isosorbide                   |
| 3   | 12.933 | 560231  | 0.06  Furan, 3-(4-methyl-5-trans-phenyl-1,3-oxazolidin-2-yl)- |
| 4   | 13.854 | 14658815 | 14.62 1,3,2,5-Dimethylene-4-rhamnitol |
| 5   | 14.235 | 38707499 | 3.86  D-Glycero-D-galacto-heptose  |
| 6   | 14.608 | 21753970 | 2.17  Lauric acid methyl ester    |
| 7   | 15.280 | 11348450 | 1.13  Tridecanoic acid            |
| 8   | 15.501 | 4033264  | 0.40  Megastigmatrieneone         |
| 9   | 16.125 | 55318375 | 5.52  2-Deoxy-d-galactose         |
| 10  | 16.648 | 30284734 | 3.02  D-Glucitol, 2,5-anhydro-1-O-octyl-    |
| 11  | 17.070 | 6971059  | 0.70  2-Hydroxyhexadecyl butanoate |
| 12  | 17.456 | 8567378  | 0.85  Myristic acid methyl ester  |
| 13  | 17.725 | 2962331  | 0.30  9-(3,3-Dimethyloxiran-2-yl)-2,7-dimethylnona-2,6-dien-1-ol |
| 14  | 18.153 | 4592151  | 0.46  6-Hydroxy-4,4,7a-trimethyl-5,6,7,7a-tetrahydrobenzofuran-2(4H)-one |
| 15  | 18.416 | 10513893 | 1.05  Benzenesulfonamide, N-butyl- |
| 16  | 18.952 | 13881898 | 1.38  Neophytadiene               |
| 17  | 19.037 | 7043508  | 0.70  2-Pentadecanone, 6,10,14-trimethyl-   |
| 18  | 19.269 | 4920685  | 0.49  [1,1′-Bicyclopropyl]-2-octanoic acid, 2′-hexyl-, methyl ester |
| 19  | 19.500 | 7077957  | 0.71  3,7,11,15-Tetramethyl-2-hexadec-1-ol |
| 20  | 20.027 | 17060966 | 17.02  Hexadecanoic acid, methyl ester |
| 21  | 20.515 | 51495640 | 5.14  Palmitic acid               |
| 22  | 21.239 | 2788195  | 0.28  Cyclohexanebutanoic acid     |
| 23  | 21.920 | 3248248  | 0.32  Methyl trans-4-[2-nonylcyclopentyl] butanoate |
| 24  | 22.033 | 9779382  | 0.98  cis-11,14-Eicosadienoic acid, methyl ester |
| 25  | 22.105 | 34030060 | 3.39  Linolenic acid methyl ester  |
| 26  | 22.172 | 3962623  | 0.40  9-Octadecenoic acid (Z), methyl ester |
| 27  | 22.247 | 21622122 | 21.55  Phytol, acetate             |
| 28  | 22.386 | 56126780 | 5.60  Methyl stearate             |
| 29  | 22.876 | 3903372  | 0.39  Benzyl beta-D-glucoside      |
| 30  | 24.565 | 3540084  | 0.35  Methyl 18-methyl nonadecanoate |
| 31  | 24.857 | 1679444  | 0.17  4,8,12,16-Tetramethylheptadecan-4-olide |
| 32  | 25.213 | 1068195  | 0.11  3,7,11,15-Tetramethylhexadec-2-en-1-yl acetate |
| 33  | 26.211 | 1769239  | 0.18  1-Heptacosanol               |
| 34  | 27.057 | 2638778  | 0.26  Phytol palmitate            |
| 35  | 28.117 | 110457  | 0.11  9-Hexacosene                 |
| 36  | 28.430 | 1074607  | 0.11  Methyl 18-methyl nonadecanoate |
| 37  | 28.792 | 1507426  | 0.15  Neophytadiene               |
| 38  | 29.305 | 1886233  | 0.19  2,6,10,14,18-Pentamethyl-2,6,10,14,18-eicosapentaene |
| 39  | 31.856 | 19787504 | 1.97  Dl-α-Tocopherol              |
| 40  | 33.332 | 3372826  | 3.36  β-Sitosterol                 |
| 41  | 34.009 | 802361   | 0.08  Acetyl betulinaldehyde       |
Based on the GC/MS analysis, it was found that most of the identified compounds were present in both *C. hissaricus* and *C. hsingshangensis*. However, (E)-2,6-dimethoxy-4-(prop-1-en-1-yl)phenol, 1-docosanol, methyl ether, 3-hydroxy-β-damascone, cholesta-4,6-dien-3-ol, (3β)-octatriacontyl trifluoroacetate, and urs-12-en-28-al were only detected in *C. hissaricus* leaves, and 2,6,10,14,18-pentamethyl-2,6,10,14,18-eicosapentaene, benzene-sulfonamide, N-butyl-α-D-glucitol, 2,5-anhydro-1-O-octyl-α-D-glucitol, lauric acid methyl ester, and tridecanoic acid were only found in *C. hsingshangensis*.
It is worth mentioning that the pronounced bioactivity of phenols toward skin disorders is enhanced by the simultaneous presence of such components as 6-hydroxy-4,4,7a-trimethyl-5,6,7,7a-tetrahydrobenzofuran-2(4H)-one, which was reported as a strong anti-inflammatory agent [56], and which was identified in both studied Cotoneaster species.

A broad variety of active compounds in Cotoneaster species lead to overlapping activities of miscellaneous agents, hence acting on several levels of different diseases caused by oxidative stress is possible.

3.2. Biological Activity

Taking into account that both oxidation and inflammation play significant roles in disease pathogenesis, recent research has focused on the evaluation of the antioxidant and anti-inflammatory activity of plants. In our study, we examined in vitro antioxidant, anti-cyclooxygenase, anti-lipooxygenase, anti-hyaluronidase, antibacterial, and cytotoxic activities of leaves of C. hissaricus and C. hsingshangensis.

3.2.1. Antioxidant Activity

The antioxidant activity was studied on the microplate scale in cell-free systems. The Cotoneaster extracts were evaluated in a concentration ranging from 10 to 150 μg/mL. It was demonstrated that all investigated extracts exhibited moderate scavenging capacity in a concentration-dependent manner (Table 6). For comparison, the radical scavenging activity of ascorbic acid (AA; IC\(_{50} = 4.75 \pm 0.16 \mu g/mL\)), quercetin (IC\(_{50} = 2.05 \pm 0.10 \mu g/mL\)) and Trolox (IC\(_{50} = 3.68 \pm 0.09 \mu g/mL\)) were tested in the same conditions. The highest DPPH scavenging activity was shown for the ethyl acetate fraction (CHs-4) (IC\(_{50} = 2.08 \pm 0.03 \mu g/mL\)), followed by the butanol fraction (CHs-3), and the diethyl ether fraction (CHs-2) of C. hissaricus activities of leaves of C. hissaricus followed by the butanol fraction (CHs-3), and the diethyl ether fraction (CHs-2) of C. hsingshangensis (IC\(_{50} = 3.43 \pm 0.02\) and 4.15 ± 0.05 μg/mL, respectively). The weakest activity was noted for the water fraction (CHi-1; IC\(_{50} = 32.37 \pm 0.19\) μg/mL) and crude extract of C. hissaricus (CHi; IC\(_{50} = 21.73 \pm 0.13\) μg/mL).

Table 6. The IC\(_{50}\) values determined in antioxidant tests.

| Sample     | DPPH [μg/mL] | ABTS [μg/mL] | CHEL [μg/mL] |
|------------|--------------|--------------|--------------|
| CHi        |              |              |              |
| CHi-1      | 21.73 ± 0.13 | 4.14 ± 0.10  | 3.54 ± 0.15  |
| CHi-2      | 32.37 ± 0.19 | 5.48 ± 0.12  | 182.67 ± 4.19|
| CHi-3      | 8.83 ± 0.10  | 1.68 ± 0.03  | 2.94 ± 0.03  |
| CHi-4      | 13.69 ± 0.14 | 2.74 ± 0.11  | 4.04 ± 0.16  |
| CHi-5      | 5.40 ± 0.10  | 0.90 ± 0.02  | 1.19 ± 0.05  |
| CHi-6      | 10.16 ± 0.02 | 1.92 ± 0.13  | 1.73 ± 0.10  |
| CHi-7      | 18.15 ± 0.13 | 1.53 ± 0.09  | 76.50 ± 1.45  |
| CHi-8      | 4.15 ± 0.05  | 0.80 ± 0.02  | 1.10 ± 0.03  |
| CHi-9      | 3.43 ± 0.02  | 0.68 ± 0.05  | 1.01 ± 0.01  |
| CHi-10     | 2.08 ± 0.03  | 0.37 ± 0.01  | 0.50 ± 0.01  |
| quercetin  | 2.05 ± 0.10  | 3.27 ± 0.23  | 6.24 ± 0.19  |
| AA         | 4.75 ± 0.16  | 1.73 ± 0.09  | nt           |
| Trolox     | 3.68 ± 0.09  | 1.31 ± 0.05  | nt           |

Data were expressed as mean ± SD; n = 3. AA—ascorbic acid; Na\(_2\)EDTA*2H\(_2\)O—ethylenediaminetetraacetic acid, disodium dihydrate; nt—not tested; CHi—crude extract of C. hissaricus; CHs—crude extract of C. hsingshangensis; CHi-1/CHs-1—water fraction, CHi-2/CHs-2—diethyl ether fraction, CHi-3/CHs-3—butanol fraction, CHi-4/CHs-4—ethyl acetate fraction; ABTS—2,2′-azino-bis-(3-ethylbenzothiazoline-6-sulfonic acid); CHEL—metal chelating activity. Statistical analysis: a—significantly different results compared to CHi-1; b—significantly different results compared to CHi-2; c—significantly different results compared to CHi-3; d—significantly different results compared to CHi-4; e—significantly different results compared to CHs-1; f—significantly different results compared to CHs-2; g—significantly different results compared to CHs-3; h—significantly different results compared to CHs-4; j—significantly different results compared to quercetin; k—significantly different results compared to AA; l—significantly different results compared to Trolox; m—significantly different results compared to Na\(_2\)EDTA*2H\(_2\)O; One-Way ANOVA test, followed by a Tukey’s multiple comparison test, p < 0.05.
The results of other published reports seem to be difficult to compare due to the other conditions used during experiments. Nevertheless, antioxidant activity using the DPPH assay was studied for different fractions of 70% methanol extract of the leaves of three Cotoneaster species cultivated in Poland [29]. The authors found that the studied extracts and fractions possessed significant scavenging effects with IC\(_{50}\) values ranging from 3.19 \(\mu g/mL\) for the ethyl acetate fraction of \textit{C. bullatus} to 27.88 \(\mu g/mL\) for the water residue of \textit{C. integerrimus} (the IC\(_{50}\) values for positive controls—quercetin, (−)-epicatechin, chlorogenic acid, butylated hydroxyanisole, and Trolox were 1.70, 2.35, 4.60, 2.90, and 4.05 \(\mu g/mL\), respectively). Holzer et al. [26] also studied antioxidant capacity with DPPH tests of different extracts of sterile shoots, old stems, and leaves of \textit{C. melanocarpus}. They found that the water extract had IC\(_{50}\) values from 53.53 to 86.72 \(\mu g/mL\), methanolic extracts—30.91–106.41 \(\mu g/mL\), ethyl acetate extracts—74.88–200 \(\mu g/mL\), dichloromethane extracts—134.60–200 \(\mu g/mL\). Ten polyphenol compounds isolated from the leaves of \textit{C. bullatus} and \textit{C. zabelii} and crude hydroalcoholic extracts were also evaluated by the DPPH radical scavenging assay. All these compounds showed significant inhibitory activity. Importantly, (−)-epicatechin (IC\(_{50}\) = 2.35 \(\mu g/mL\)), procyanadin B2 (IC\(_{50}\) = 2.46 \(\mu g/mL\)), and procyanadin C1 (IC\(_{50}\) = 2.55 \(\mu g/mL\)) showed stronger activity compared to Trolox (IC\(_{50}\) = 4.06 ± 0.11 \(\mu g/mL\)), which was used as a positive control [36]. Significant antioxidant activities were also observed for the compounds isolated from the leafy twigs of \textit{C. racemiflora}, including racemiside (IC\(_{50}\) = 11.1 \(\mu g/mL\)), scopoletin (IC\(_{50}\) = 3.8 \(\mu g/mL\)), 7,8-dimethoxy-6-hydroxycoumarin (IC\(_{50}\) = 1.8 \(\mu g/mL\)), 3,3′,4′-tri-O-methyllellagic acid (IC\(_{50}\) = 15.1 \(\mu M\)), and cereotagloperoxide (IC\(_{50}\) = 5.9 \(\mu M\)) (IC\(_{50}\) for BHA used as positive control was 44.2 \(\mu M\)) [44].

As shown in Table 6, similar to the DPPH test, the ABTS\(^{••}\) assay revealed that the ethyl acetate fraction (CHs-4) of the leaves of \textit{C. hsingshangensis} possessed the strongest ability to scavenge free radicals (IC\(_{50}\) = 0.37 ± 0.01 \(\mu g/mL\)), followed by the butanol fraction (CHs-3; IC\(_{50}\) = 0.68 ± 0.05 \(\mu g/mL\)) and the diethyl ether fraction (CHs-2; IC\(_{50}\) = 0.80 ± 0.02 \(\mu g/mL\)) of \textit{C. hsingshangensis}. The ABTS\(^{••}\) assay was also used to test different extracts of \textit{C. nummularia}. The authors found that the greatest ABTS inhibition was caused by the methanol (IC\(_{50}\) = 0.020 mg/mL) and the diethyl ether fraction (CHs-2; IC\(_{50}\) = 0.02 mg/mL) (IC\(_{50}\) for BHA used as a positive control was 0.15 mg/mL) [57].Mahmutović-Dizdarević and co-authors [11] reported that IC\(_{50}\) values of methanolic extracts of leaves and barks of \textit{C. horizontalis}, \textit{C. integerrimus}, and \textit{C. tomentosum} ranged from 0.12 (leaves of \textit{C. tomentosum}) to 0.87 mg/mL (barks of \textit{C. tomentosum}).

Phenolic compounds can reduce oxidative stress by several mechanisms that depend on their chemical structure. One of them is the chelation of metal ions, such as iron, which plays a key role in the production of damaging oxygen species [58].

The chelating ability was determined based on measurement of the percentage of inhibition of formation of the ferrozine-Fe\(^{2+}\) complex. As reported in Table 6, the extracts and fractions from leaves of both studied Cotoneaster species possessed the capacity to interfere with the formation of iron and ferrozine complexes, which suggests their high chelating capacity and ability to capture iron ions before ferrozine. The IC\(_{50}\) values of most of these extracts showed higher chelating activity than the positive control—\(\text{Na}_2\text{EDTA}·2\text{H}_2\text{O}\) (IC\(_{50}\) = 4.15 ± 0.10 \(\mu g/mL\)). Among the investigated extracts, the best activity was noticed in the ethyl acetate fraction of \textit{C. hsingshangensis} (IC\(_{50}\) = 0.50 \(\mu g/mL\)). Zengin and co-authors [57] also evaluated the metal chelating capacity of different extracts of aerial parts of \textit{C. nummularia}. They found that the highest chelating activity was expressed by the ethyl acetate extract (IC\(_{50}\) = 0.25 mg EDTA/g of extract). Among the ethanol extracts of 34 species belonging to Rosaceae, Cotoneaster meyeri, \textit{C. morulus}, and \textit{C. nummularia} caused moderate metal-chelating effects (5.91, 21.48, and 26.19 chelation%, respectively) [34].

3.2.2. Enzyme Inhibitory Activity

Lipoxygenases (LOXs) are present in the human body and play a crucial role in the stimulation of inflammatory reactions. Exaggerated quantities of reactive oxygen species
can induce inflammation that stimulates the release of cytokines and then the activation of lipoxygenases. They are connected with the spread of many diseases, and their inhibition is viewed as a relevant step in their prevention [59]. The lipoxygenase inhibitor screening assay kit is a popular method for lipoxygenase detection, which estimates the presence of hydroperoxides (4-hydroperoxy cis-trans 1,3-conjugated pentadienyl moiety within the unsaturated fatty acid) at various positions (5, 12-, 15-), which are produced in the lipoxygenation reaction using a purified lipoxygenase.

Sinha et al. [60] noted that peroxisome proliferator activated receptor (PRAR) ligands induce lipogenesis in cultured human sebocytes; hence, 5-lipoxygenase inhibitors demonstrate the ability to decline acne lesions, due to the fact that they possess the capability to reduce lipogenesis. The presence of PPARα receptors within sebocytes has been observed in peroxisomes, mitochondria, and microsomes.

The results of the inhibition of lipoxygenase are shown in Table 7. The diethyl ether fraction (CHs-2) and ethyl acetate fraction (CHs-4) of *C. hsingshangensis* showed a considerable ability to inhibit lipoxygenase activity (IC$_{50} = 4.15$, and 5.72 µg/mL, respectively), while the water fraction of *C. hissaricus* showed the lowest activity (IC$_{50} = 129.46$ µg/mL). Both CHs-2 and CHs-4 exhibited significantly higher inhibitory activity than that of nordihydroguaiaretic acid (NDGA) used as a positive standard (IC$_{50} = 5.89$ µg/mL). In a recent study of *Cotoneaster* species in regard to their capacity to inhibit lipoxygenase, it was found that different extracts of the leaves of *C. bullatus*, *C. integerrimus*, and *C. zabelii* have moderate activity with IC$_{50}$ values in the range of 95.19 to 684.84 µg/mL. The best LOX inhibition was achieved by the butanol fraction of *C. bullatus* (IC$_{50} = 95.19$ µg/mL) [29]. The hydromethanolic extracts of fruits of nine *Cotoneaster* species cultivated in Poland showed the strongest inhibition of LOX, with IC$_{50}$ values in the range of 62.54 (C. *zabelii*) to 165.76 (C. *nanshan*) µg/mL [28].

### Table 7. Anti-lipoxygenase, anti-hyaluronidase, and anti-cyclooxygenase activities of the leaves of *C. hissaricus* and *C. hsingshangensis*.

| Sample   | Lipoxygenase Inhibition | Hyaluronidase Inhibition | IC$_{50}$ [µg/mL] |
|----------|-------------------------|-------------------------|-------------------|
| CHi      | 28.73 ± 2.01 c,f,g,h,i,j | 15.09 ± 0.61 e,g,h,i,j  | 24.77 ± 0.35 e,f,g,h,i,j |
| CHi-1    | 129.46 ± 6.35 a,i,l     | 24.95 ± 0.52 a,b,c,e,f,g,h,i,j | 62.54 ± 1.02 a,b,c,e,f,g,h,i,j |
| CHi-2    | 19.77 ± 1.56 b,i,l      | 13.19 ± 0.08 a,b,c,e,f,g,h,i,j | 11.15 ± 0.42 a,b,c,e,f,g,h,i,j |
| CHi-3    | 45.69 ± 4.21 a,b,c,i,l  | 10.23 ± 0.23 a,b,c,e,f,g,h,i,j | 46.71 ± 0.69 a,b,c,e,f,g,h,i,j |
| CHi-4    | 97.12 ± 3.86 a,b,c,d,i,l| 19.80 ± 0.51 a,b,c,e,f,g,h,i,j | 45.35 ± 0.47 a,b,c,e,f,g,h,i,j |
| CHs      | 11.06 ± 1.14 i          | 6.82 ± 0.15 k           | 13.02 ± 0.17 i     |
| CHs-1    | 72.15 ± 1.57 c,i        | 14.76 ± 0.19 e          | 34.12 ± 0.28 c,i   |
| CHs-2    | 4.15 ± 0.31 c,f         | 1.17 ± 0.02 e,f         | 6.39 ± 0.04 e,f    |
| CHs-3    | 11.56 ± 0.98 t,g         | 7.41 ± 0.05 e,f,g,k     | 15.56 ± 0.13 e,f,g,i,j |
| CHs-4    | 5.72 ± 0.26 c,f,h       | 1.89 ± 0.05 e,f,g,h,k   | 9.54 ± 0.04 e,f,g,i,j |
| IND      | nt                      | nt                      | 4.34 ± 0.05        |
| EGCG     | nt                      | nt                      | 6.25 ± 0.02        |
| NDGA     | 5.89 ± 0.15             | nt                      | 3.82 ± 0.09        |

CHi—crude extract of *C. hissaricus*, CHs—crude extract of *C. hsingshangensis*, CHi-1/CHs-1—water fraction, CHi-2/CHs-2—diethyl ether fraction, CHi-3/CHs-3—butanol fraction, CHi-4/CHs-4—ethyl acetate fraction; EGCG—epigallocatechin gallate, NDGA—nordihydroguaiaretic acid; IND—Indomethacin. Statistical analysis: a—significantly different results compared to CHi; b—significantly different results compared to CHi-1; c—significantly different results compared to CHi-2; d—significantly different results compared to CHi-3; e—significantly different results compared to CHs; f—significantly different results compared to CHs-1; g—significantly different results compared to CHs-2; h—significantly different results compared to CHs-3; i—significantly different results compared to CHs-4; j—significantly different results compared to IND; k—significantly different results compared to EGCG; l—significantly different results compared to NDGA; One-Way ANOVA test, followed by a Tukey’s multiple comparison test, p < 0.05.
Skin is especially sensitive to reactive oxygen species because it is exposed to oxidative stress from both endogenous and exogenous sources. Although oxidative stress is a key factor in this process, hyaluronic acid also plays a notable role. Its integrity inside the dermal matrix is substantial for cell integrity and proliferation. Under oxidative stress, hyaluronidase, which is responsible for hyaluronic acid depolymerization, is overactivated and breaks down this anionic glycosaminoglycan, carrying on to the destruction of the proteoglycan system. This results in the deregulation of skin homeostasis and increases inflammatory and allergic conditions [61,62].

In our study, the ethyl acetate fraction of \textit{C. hsingshangensis} exhibited the best hyaluronidase inhibition activity (IC$_{50}$ = 1.89 µg/mL) compared to the other extracts (Table 7). Most importantly, such activity was even better than the positive standard—EGCG (IC$_{50}$ = 6.25 µg/mL). In turn, the IC$_{50}$ values for the crude extracts of \textit{C. hissaricus} and \textit{C. hsingshangensis} were 15.09 ± 0.61, and 6.82 ± 0.15 µg/mL, respectively. Previous studies have shown that methanol extracts of the leaves of \textit{C. bullatus}, \textit{C. integerrimus}, and \textit{C. zabelii} reduced the activity of hyaluronidase in a concentration-dependent manner. The most active were the butanol fraction of \textit{C. bullatus} and \textit{C. zabelii} (IC$_{50}$ = 2.81, and 6.08 µg/mL, respectively) and they had inhibition activity better than indomethacin used as a positive control (IC$_{50}$ = 8.61 µg/mL) [29]. Moreover, methanol extracts of fruits of \textit{C. bullatus}, \textit{C. dielsianus}, \textit{C. divaricatus}, \textit{C. hjelmqvistii}, \textit{C. horizontalis}, \textit{C. lucidus}, \textit{C. nanshan}, \textit{C. splendens}, and \textit{C. zabelii} showed moderate inhibition of LOX with IC$_{50}$ values in the range of 25.65 (\textit{C. lucidus}) to 45.64 µg/mL (\textit{C. nanshan}) [28].

Cyclooxygenases catalyze two reactions, the first being a cyclooxygenase function consisting of the addition of molecular oxygen to arachidonic acid to form prostaglandin G$_2$ (PGG$_2$). The second is the conversion of PGG$_2$ to PGH$_2$ by a peroxidase function. Therefore, this enzyme performs the initial reaction in the arachidonic acid metabolic cascade, carrying on to the formation of pro-inflammatory, i.e., prostaglandins, which regulate smooth muscle contractility, platelet aggregation, and mediate pain. Cyclooxygenase constitutive (COX-1) is responsible for the maintenance of physiological prostanoïd biosynthesis, and COX-2 (an inducible isoform) is connected to inflammatory cell types and tissues [63].

To determine the potential anti-inflammatory properties of \textit{Cotoneaster} leaves, we also estimated the ability of the extracts and fractions to inhibit the conversion of arachidonic acid to PGH$_2$ by ovine COX-1 and human recombinant COX-2 using a COX inhibitor screening assay kit (Cayman Chemical, MI, USA). As shown in Table 7, most of the studied extracts and fractions showed good activity against COX-1 and COX-2. The most active against COX-1 were CHs-2 (IC$_{50}$ = 6.39 µg/mL) and CHs-4 (IC$_{50}$ = 9.54 µg/mL), followed by CHi-2 (IC$_{50}$ = 11.15 µg/mL) and CHs (IC$_{50}$ = 13.02 µg/mL), while the weakest was fraction CHi-1 (IC$_{50}$ = 62.54 µg/mL). Except for the water residual of \textit{C. hissaricus}—CHi-1 (IC$_{50}$ = 100.36 µg/mL) and ethyl acetate fraction of the \textit{C. hissaricus}—CHi-4 (IC$_{50}$ = 81.69 µg/mL), all extracts and fractions of both \textit{Cotoneaster} species showed good activity against COX-2 (IC$_{50}$ = 5.09–57.59 µg/mL).

3.2.3. Antibacterial Activity

Diffusion Test in Solid Medium

The antibacterial activity of the \textit{Cotoneaster} extracts and fractions was assessed by diffusion test in a solid medium and measuring the zones of bacterial growth inhibition. The larger the zone around the applied samples, the more active the extract/fraction is.

The data show (Figure 7, Table S1) that the \textit{Cotoneaster} fractions, not the starting crude extracts, possessed the strongest antimicrobial properties. The largest zones of growth inhibition of Gram-positive microaerobic bacteria were produced by CHs-2 fraction (23 mm–21 mm), then CHs-4 (19 mm–16 mm), CHs-1 (15 mm–14 mm), and CHs-3 fractions (14 mm–11 mm). These fractions against Gram-positive aerobic strains showed weaker activity (in the range 19 mm–6 mm) than against microaerobic bacteria. Crude \textit{Cotoneaster} extracts CHi and CHs showed moderate activity against all tested Gram-positive strains (12 mm–6 mm). None of the samples were active against Gram-negative bacteria. Previous
studies have reported the resistance of both Gram-positive and Gram-negative bacteria to the leaf and bark of *C. integrerrimus*, *C. tomentosus*, and *C. horizontalis* methanolic extracts. Mahmutović-Dizdarević and co-authors [11] found that these extracts have significant antimicrobial activity against *Salmonella enterica, Pseudomonas aeruginosa, Escherichia coli*, Extended Spectrum Beta-Lactamase producing *E. coli* or ESBL *E. coli*, *Enterococcus faecalis*, *Staphylococcus aureus*, Methicillin-resistant *Staphylococcus aureus* or MRSA, and *Bacillus subtilis*. In a further study, the antibacterial activity of the ethanolic extract of roots of *C. acuminatus* was also investigated against *Bacillus pumilus, B. subtilis, E. coli, Micrococcus glutamicus, S. aureus*, *Proteus vulgaris*, and *P. aeruginosa*. The greatest effect was found for 100 μg/mL extract, with growth inhibition zones 10–18 mm [64]. Using the broth microdilution method, the antibacterial activity of the water, methanol, and ethyl acetate extracts of *C. nummularia* was evaluated against *E. coli, Bacillus cereus, P. aeruginosa*, methicillin sensitive *S. aureus* (MSSA), *Klebsiella pneumoniae*, *Salmonella enteritidis*, *S. pneumoniae*, *Sarcina lutea*, *Enterococcus faecalis*, methicillin resistant *S. aureus* (MRSA), and strains of methicillin resistant *S. aureus* isolated from clinical samples. The authors found that *E. faecalis* was the most sensitive bacteria, and *B. cereus, K. pneumoniae*, and *S. enteritidis* were the most resistant bacteria against all extracts except for the ethyl acetate extract [57].

![Zones of bacterial growth inhibition](image)

**Figure 7.** Zones of bacterial growth inhibition of the *Cotoneaster* extracts.

To the best of our knowledge, this study represents the first report of the antibacterial activity against *C. acnes, S. epidermidis*, and *C. granulosum* strains, which can be responsible for skin diseases of *Cotoneaster* species.

Results of MIC and MBC Determination

In order to determine the MIC values of the selected active fractions of crude *C. hsingshangensis* extract (CHi, CHs, CHs-1-CHs-4), an assay was performed using the double dilution method in a 96-well plate. After the plates were incubated under the appropriate conditions for the given strain, they were analyzed in an automatic plate reader (at 600 nm) against the control. The concentration at which no bacterial growth was observed was taken as the MIC. Next, the MBC value was determined by dispensing 10 μL of the solutions taken from the wells of 96-well plates in which no bacterial growth was observed.

The data contained in Table 8 confirm the results obtained in the earlier diffusion assay; namely, the tested fractions are the most active against the acne strains of *Cutibacterium* spp.
Table 8. Minimum inhibitory concentration (MIC) and minimum bactericidal concentration of the fractions of *C. hsingshangensis* crude extract.

| Sample     | S. aureus ATCC 25923 | S. epidermidis ATCC 12228 | C. acnes PCM 2400 | C. acnes PCM 2334 | C. acnes ATCC 11827 | C. granulosus PCM 2462 |
|------------|----------------------|---------------------------|-------------------|-------------------|---------------------|-----------------------|
|            | MIC [µg/mL] | MBC [µg/mL] | MIC [µg/mL] | MBC [µg/mL] | MIC [µg/mL] | MBC [µg/mL] | MIC [µg/mL] | MBC [µg/mL] | MIC [µg/mL] | MBC [µg/mL] | MIC [µg/mL] | MBC [µg/mL] |
| CHi        | >1000     >4       | >1000   >4       | >1000   >4       | >1000   >4       | >1000   >4       | >1000   >4       | >1000   >4       | >1000   >4       |
| CHs        | >1000     >4       | >1000   >4       | >1000   >4       | >1000   >4       | >1000   >4       | >1000   >4       | >1000   >4       | >1000   >4       |
| CHs-1      | >1000     >4       | 1000    >4       | 500     >4       | 1000    >4       | 1000    >4       | 1000    >4       | 500     >4       | >1000   >4       |
| CHs-2      | 500       >4       | 1000    >4       | 31.25   8       | 62.5    8       | 125     8       | 62.5    8       | >250     >4       | >250     >4       |
| CHs-3      | >1000     >4       | >1000   >4       | 1000    >4       | 500     >4       | 1000    >4       | 1000    >4       | >250     >4       | >250     >4       |
| CHs-4      | 1000      >4       | >1000   >4       | 250     >4       | 250     >4       | 125     >8       | 250     >4       | >250     >4       | >250     >4       |

CHs-1—water fraction of *C. hsingshangensis*, CHs-2—diethyl ether fraction of *C. hsingshangensis*, CHs-3—butanol fraction of *C. hsingshangensis*, CHs-4—ethyl acetate fraction of *C. hsingshangensis*.

The most favorable, i.e., the lowest MIC values against these bacteria, were obtained by fractions CHs-2 (MIC 31.25–125 µg/mL), CHs-4 (125–250 µg/mL), CHs-1, and CHs-3 (both 500–1000 µg/mL). Next, the MBC—as the lowest concentration of an antibacterial sample required to kill the bacterium—can be assayed based on an MIC test by subculturing samples on agar plates. Thus, the clear medium from MIC plates was spread on fresh agar to check its sterility. The concentration of the sample that did not produce colonies was considered the MBC power. According to the ratio MBC/MIC, we visualized and appreciated antibacterial activity. If the ratio MBC/MIC $\leq 4$, the effect was considered bactericidal, but if the ratio MBC/MIC $> 4$, the effect was defined as bacteriostatic. Data of MBC/MIC ratio displayed in Table 8 show that none of *C. hsingshangensis* fractions showed any bactericidal activity in relation to the acne bacteria. The most promising power against microaerobic Gram-positive strains was displayed for CHs-2 (diethyl ether fraction of *C. hsingshangensis*). The remaining extract power (CHs-1, CHs-3, CHs-4) was not measurable.

### 3.2.4. Cytotoxic Activity

The obtained data revealed that all *Cotoneaster* extracts possessed low cytotoxicity toward normal human fibroblasts (Figure 8), as it was not possible to determine their CC$_{50}$ values at tested concentrations (1.95–1000 µg/mL). The CC$_{50}$ value denotes the extract concentration that inhibits BJ viability to 50%. Among the tested *Cotoneaster* extracts, CHs-3 exhibited the lowest ability to inhibit the viability of BJ cells because at the highest tested concentration (1000 µg/mL), the cell viability was 77.06 ± 1.57%. For comparison, fibroblast viability treated with CHi, CHs, CHs-1, and CHs-4 at the same concentration was 60.60 ± 1.47%, 53.18 ± 1.84%, 68.28 ± 5.55%, and 63.80 ± 4.39%, respectively. In the case of CHs-2, cell viability after incubation at concentrations of 250 µg/mL, 500 µg/mL, and 1000 µg/mL was 54.40 ± 3.56%, 43.35 ± 2.02%, and 45.89 ± 4.57%, respectively. Thus, these results may suggest that the CC$_{50}$ value for this substance should be detected in the range 250–500 µg/mL. As a consequence, it was demonstrated that CHs-2 possessed the highest ability to inhibit BJ cell viability compared to other substances.

### 3.3. Multivariate Analysis of the Results

To perform a holistic view of the results, we used a chemometric multivariate approach. It allows us to examine some independent trends in changes among the investigated properties. They were investigated using hierarchical cluster analysis (HCA, Figure 9) and principal component analysis (PCA, Figure 10).
The analysis was done on a column-wise scaled matrix, as each parameter is expressed with different units. Additionally, two of the parameters, DPPH and ABTS, were expressed as negative values in this matrix. This was done to convert the strong correlation between them and TPC, TPAC, and TFC from negative to positive. This places the loading vectors in the same direction and allows us to see everything visually in more detail.

It can be clearly seen that the parameters form two distinct groups, expressed as two separate groups of loading arrows and two main branches of the dendrogram. The first group contains antioxidant parameters TPC, TPAC, and TFC, together with DPPH and ABTS. The other parameters form the second group.

The increase or decrease of all investigated parameters in opposite directions is responsible for 75% of total variance and is modeled as PC1. PC2 represents an intercorrelated increase or decrease of all investigated parameters and is responsible for 14% of the variability. Further PCs do not contain any interpretable information (results not shown).

Figure 8. Cytotoxic effect of Cotoneaster extracts on human normal fibroblasts. (BJ cell line, ATCC CRL-2522TM). The cell viability was assessed after 24-h incubation using the MTT assay. Asterisk (*) denotes significantly different data ($p < 0.05$, unpaired t-test) compared to the control, namely culture medium without substances—0 μg/mL.
Figure 9. The heatmap analysis of the scaled matrix, with dendrograms based on Euclidean distance. TPC—Total phenolic content; TPCA—Total phenolic acids content; TFC—Total flavonoid content; CHEL—Metal chelating activity; LPO—Lipoxygenase inhibition; HYAL—Hyaluronidase inhibition.

Figure 10. Principal component analysis loadings of the investigated dataset: PC1 vs. PC2 plot. TPC—Total phenolic content; TPCA—Total phenolic acids content; TFC—Total flavonoid content; CHEL—Metal chelating activity; LPO—Lipoxygenase inhibition; HYAL—Hyaluronidase inhibition.
4. Conclusions

The management of acne vulgaris should be “multidirectional”. Therapeutics should reduce sebum production, as well as follicular hyperkeratinization of the epidermal cells. For this reason, they should primarily exhibit antioxidant, anti-inflammatory, and antimicrobial activities, without cytotoxic effects.

Our analysis has been designed due to missing data in the available literature, namely considerable gaps in the knowledge considering C. hissaricus and C. hsingshangensis are observed. The unique chemical composition of the representatives of the Rosaceae family makes it possible to obtain a wide range of pharmaceutical, medicinal, and cosmetic products. Thus, Cotoneaster species seem to be promising candidates for further research. Chemical drugs have limitations because of their high toxic activity, as well as their ability to induce adverse side effects. Thus, a growing interest in the use of herbal medicines for the management of acne and other diseases is increasingly recognized.

Among approximately 2500 species belonging to the family Rosaceae, in vitro anti-acne research has been carried out only with a small number of them, namely Rosa damascena [65], Rosa multiflora [66], Prunus jamasakura [67], and apple polyphenols [54]. Interestingly, there is a lack of research on other plants from the Rosaceae family for anti-acne activity and the effect of phytochemicals on the skin. Simultaneously, indisputable evidence that phenolic compounds exhibit the desired effects and contribute to alleviate skin disorders can be found in the available literature [51,68]. Furthermore, clindamycin and kaempferol in combination with quercetin possess more beneficial effects than other formulations [69].

In our in vitro research, we characterized composition and evaluated the biological properties of extracts from C. hissaricus and C. hsingshangensis. Thus, we identified the main compounds present in extracts, as well as determined the antioxidant, anti-inflammatory, antimicrobial, and cytotoxic properties of such extracts.

To sum up our results, it seems to be clear that comprehensive and well-designed future research on phenolic compounds from Cotoneaster in greater detail will constitute significant importance in pharmacy and medicine. Among the studied extracts, the diethyl ether fraction of C. hsingshangensis (CHs-2) exhibited great ability to scavenge free radicals and good capacity to inhibit cyclooxygenase-1, cyclooxygenase-2, lipoxygenase, and hyaluronidase. Moreover, it had the most promising power against microaerobic Gram-positive strains, and importantly, it was non-toxic toward normal skin fibroblasts. Taking into account the value of the calculated therapeutic index (>10), it is worth noting that CHs-2 can be subjected to in vivo study and constitutes a promising anti-acne agent.

To the best of our knowledge, there is no reported study on the chemical composition and skin-related properties of the examined species.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/cells11030367/s1, Table S1: Zones of bacterial growth inhibition of the Cotoneaster extracts and fractions.

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