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REVIEW ARTICLE

Antioxidant potential of lichen species and their secondary metabolites. A systematic review

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Abstract

Context: Pharmacological interest of lichens lies in their capacity to produce bioactive secondary metabolites, being most of them phenolic compounds with reactive hydroxyl groups that confer antioxidant potential through various mechanisms. Increasing incidence and impact of oxidative stress-related diseases (i.e., neurodegenerative disorders) has encouraged the search of new pharmacological strategies to face them. Lichens appear to be a promising source of phenolic compounds in the discovery of natural products exerting antioxidant activity.

Objective: The present review thoroughly discusses the available knowledge on antioxidant properties of lichens, including both *in vitro* and *in vivo* studies and the parameters assessed so far on lichen constituents.

Methods: Literature survey was performed by using as main databases PubMed, Google Scholar, Scopus, Science Direct, and Recent Literature on Lichens. We reviewed 98 highlighted research articles without date restriction.

Results: Current report collects data related to antioxidant activities of more than 75 lichen species (from 18 botanical families) and 65 isolated metabolites. Much information comes from *in vitro* investigations, such as chemical assays evaluating radical scavenging properties, lipid peroxidation inhibition, and reducing power of lichen species and compounds; similarly, research on cellular substrates and animal models generally measures antioxidant enzymes levels and other antioxidant markers, such as glutathione levels or tissue peroxidation.

Conclusion: Since consistent evidence demonstrated the contribution of oxidative stress on the development and progression of several human diseases, reviewed data suggest that some lichen compounds are worthy of further investigation and better understanding of their antioxidant and neuroprotective potentials.

Keywords

Antioxidants, neurodegenerative diseases, lichens, oxidative stress, scavenging properties

History

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Introduction

A lichen is a stable, ecologically obligate, self-supporting mutualism between an exhabitant fungus (the mycobiont) and one or more extracellularly located inhabitants, which can be either unicellular or filamentous photoautotrophic partners (the photobiont: alga or cyanobacterium) (Hawksworth & Honegger, 1994). Lichens have been used with medicinal purposes since the ancient times. For instance, *Usnea barbata* (L.) Weber ex F.H. Wigg (Parmeliaceae) and other *Usnea* species were used to treat hair-related diseases, *Lobaria pulmonaria* (L.) Hoffm. (Lobariaceae) and *Parmelia sulcata* Taylor (Parmeliaceae) for pulmonary and cranial diseases, respectively, yellow-orange colored *Xanthoria parietina* (L.) Th. Fr. (Teloschistaceae) for jaundice, *Peltigera aphthosa* (L.) Willd. (Peltigeraceae) for aphta, *Parmelia saxatilis* (L.) Ach.

(Parmeliaceae) for epilepsy, etc. (Brodo et al., 2001; Malhotra et al., 2008).

Pharmaceutical importance of lichens lies in their capacity to produce a great variety of secondary metabolites, many of which only appear in these lichenised fungi. Phenolic compounds are the most relevant secondary metabolites of lichen samples, and the best studied metabolites can be principally classified as depsides, depsidones, dibenzofurans, and pulvinic acid derivatives (Huneck, 1999). Chemical structures of some representative compounds of each group are shown in Figure 1. Systematic study of pharmacological properties of lichen compounds has recently started and they have attracted much attention in recent investigations because of their antiviral, antibiotic, antitumor, allergenic, and plant growth inhibitory activities (Dias & Urban, 2009; Einarsdóttir et al., 2010; Esimone et al., 2007; Honda & Vilegas, 1999; Nishitoba et al., 1987).

In the last years, there is an increasing interest in new bioactive natural products for the prevention and treatment of various human diseases, with remarkable attention to neurodegenerative diseases and compounds exerting

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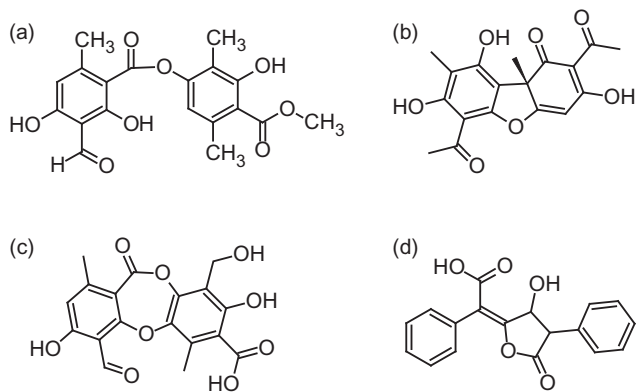


Figure 1. Chemical structures of depside atranorin (a), dibezofuran usnic acid (b), depsidone protocetraric acid (c), and pulvinic acid (d).

neuroprotective potential and oxidative stress reversion (Gonzalez-Burgos et al., 2013; Wang et al., 2013). This is due to the fact that compounds of natural origin normally have beneficial effects on the human organism with lower incidence of unwanted effects. Regarding this point, lichens are the subject of many research teams (Karakus et al., 2009; Kosanić et al., 2012b).

The present review aims to summarize and discuss the available information about the antioxidant properties of lichens and their isolated secondary metabolites in order to facilitate and guide future research on these natural products.

Oxidative stress

Free radicals (including reactive oxygen species, such as the hydroxyl radical, superoxide anion, hydrogen peroxide, and reactive nitrogen species, such as nitric oxide) play important roles in many chemical processes of the cells under physiological conditions, but they are also associated with pathology and cell damage. Oxidative stress is defined as an imbalance between biochemical processes leading to the production of reactive oxygen species (ROS) and those responsible for the removal of ROS, the antioxidant cascade. In that situation, free radicals will be able to attack nucleic acids and proteins, as well as unsaturated fatty acids in the cell membrane; several human chronic diseases (such as neurodegenerative diseases) are related to this problem (Molnár & Farkas, 2010; Sayre et al., 2008).

On the contrary, antioxidant agents inhibit and prevent those ROS that can cause degenerative diseases. Since many synthetic antioxidants have shown toxic and/or mutagenic effects (Grice, 1986; Wichi, 1988), the scientific attention shifted towards the discovery of naturally occurring antioxidants. With this regard, numerous plant constituents have been shown to exert antioxidant activity, being flavonoids and other phenolic compounds such as hydroxycinnamic derivatives, catechins, theaflavins, curcumins and terpenoids remarkable among them (Gonzalez-Burgos et al., 2012; Sundararajan et al., 2006). They are mostly phenolic compounds containing reactive hydroxyl groups and their antioxidant properties might be based on their ability to scavenge ROS, chelate metal ions (i.e., iron and copper),

stabilize unpaired electrons, and modulate the endogenous enzymatic and non-enzymatic antioxidant defense systems.

There is unquestionable evidence for some participation of oxidative stress in all neurodegenerative diseases (such Alzheimer's, Parkinson's and Huntington's diseases or amyotrophic lateral sclerosis among them). It therefore suggests possibilities of intervention into etiology or disease progression through individual or combined use of antioxidants capable to enhance endogenous enzymatic or non-enzymatic defense processes (Sayre et al., 2008). Thus, since natural antioxidants are preferred over synthetic ones and there is solid basis in thinking of a potential neuroprotective activity for phenolic compounds, investigation of the antioxidant potential of lichen metabolites becomes an interesting strategy for the prevention or treatment of various oxidative stress-related diseases.

Detailed mechanisms and pathways involved in the antioxidant activity of lichens still need further investigation. But classification of these issues known so far could help to understand the pharmacology of lichens secondary metabolites, and their possibilities in the treatment of neurodegenerative disorders, thus promoting the development of neuroprotective natural products.

Literature search

Current report is intended to discuss past and current research on antioxidant properties of lichens and their secondary metabolites. With this aim, an extensive review of scientific literature was carried out by considering all highlighted research articles and other reviews on the issue, without date or language restriction. Five main databases (PubMed, Google Scholar, Scopus, Science Direct, and Recent Literature on Lichens) were used as information sources through the inclusion of the search terms “lichens”, “lichen metabolites”, “antioxidant activities”, “oxidative stress”, and their combinations. As a result, a total of 98 bibliographic references are included in the present work.

Antioxidant properties of lichens

Some lichen extracts and metabolites have already been reported for antioxidant properties due to their phenolic content; for instance, the antioxidant activities of some depsides and depsidones isolated from several lichen species have been demonstrated (de Paz et al., 2010; Hidalgo et al., 1994; Jayaprakasha & Rao, 2000), as well as the *in vitro* properties of some lichen extracts (Gülçin et al., 2002; Stojanović et al., 2010). Even so, both studies on intracellular ROS modulation by lichen metabolites/extracts and their *in vivo* effects have been recently started.

Table 1 actually includes the antioxidant activity as revealed by *in vitro* assays of more than 75 species divided into 18 botanical families, among which Parmeliaceae family is the best studied, as it is one of the most rife and widespread. Some of the reflected studies are macrostudies in which authors evaluated the same antioxidant parameter in numerous species. Similarly, in Table 2, we gather all available data related to *in vitro* antioxidant activity of isolated lichen metabolites, referring to more than 65 compounds. In general, antioxidant activity has been mainly evaluated based on some

Table 1. Antioxidant activities of lichen extracts.

Lichen species	Solvent	LP inhibition (IC ₅₀ or %INH)	DPPH (IC ₅₀ or %INH)	SO [·] RSA (IC ₅₀ or %INH)	Reducing power (A ₇₀₀ or IC ₅₀)	Phenolic/flavonoid content	Isolated compounds/ composition	Reference
<i>Arthoniaceae</i> Reichenb. ex Reichenb. <i>Arithothelium avastii</i> Pawl. & Makhija	ME	2.1–68.4% 15.7 µg/ml	2.6–68.4% 13.2 µg/ml	Not studied (NS)	NS			Vemra et al. (2008b)
<i>Catillariaceae</i> Hafellner. <i>Toninia candida</i> (Weber) Th.Fr.	ChE, ME, PE	ChE: 41.7 µg/ml ME: 46.5 µg/ml PE: 21.5 µg/ml NS	ChE: 48.9 µg/ml ME: 51.5 µg/ml PE: 50.1 µg/ml 115.8 µg/ml	NS	ChE: 56.7 µg/ml ME: 78.5 µg/ml PE: 51.5 µg/ml 0.055 (500 µg/ml)		Noristic acid, stictic acid, usnic acid, protocetraric acid, atranorin	Manojlović et al. (2012b)
	AcE						Noristic acid, stictic acid, protocetraric acid, usnic acid, atranorin	Ranković et al. (2012)
<i>Cladoniaceae</i> Zenker. <i>Cladonia amaraeotacea</i> (Flörke) Schaefer	AcE, DmE, EE, ME	AcE: 80.3% DmE, EE, ME: numeric value not reported (NVNR)	NVNR	NS			Atranorin, usnic acid	Singh et al. (2011)
<i>Cladonia ciliata</i> Ahti & L. Xavier	EE 70%	NS	50.2% 69.3 µg/ml	NS				
<i>Cladonia digitata</i> (L.) Hofm.	ME	9.8%	NS	NS	0.085			Silva et al. (2010)
<i>Cladonia fimbriata</i> (L.) Fr.	ME	20.4%	NS	NS	0.114			Ranković et al. (2010b)
<i>Cladonia foliacea</i> (Huds.) Wild.	ME	NS	>1000 µg/ml	NS				Ranković et al. (2010b)
<i>Cladonia furcata</i> (Hudson) Schrad.	AcE, ME	NVNR	AcE: 4/71.3 µg/ml ME: >1000 µg/ml NVNR	NVNR	78.1 mg GA/g (28.2 mg Rutin/g)			Mitrović et al. (2011)
	AcE, ME, WE	NS	ME > AcE > WE	ME > AcE > WE	NS			Luo et al. (2009)
	AcE	NS	44.8%	23.9%	0.051	12.9 µg PE/mg (10.6 µg RE/mg)		Kosančić et al. (2011)
<i>Cladonia mediterranea</i> P.A. Duivinc. & Abbayes	AcE, DmE, EE, ME	AcE: 73.7% DmE, EE, ME: NVNR	NVNR	NS		AcE: 0.06 mg/g DmE: 0.001 mg/g	Usnic acid	Ranković et al. (2011)
<i>Cladonia rangiformis</i> Hofm	ChE, ME, WE	ChE: 48.3% ME: 1.6% WE: -11.5%	NS	NS	ChE: 0.083 ME: 0.115 WE: 0.034	ME: 0.009 mg/g ChE: 0.048 µg GA/ml ME: 0.010 µg GA/ml WE: 0.004 µg GA/ml	NS	Singh et al. (2011)
<i>Graphidaceae</i> Dumort. <i>Graphis guimaraiana</i> Vain.	ME 10%	NS						Yıldız et al. (2007)
<i>Graphis nakanishiana</i> Pawl. & C.R. Kulk.	ME 10%	NS						
<i>Graphis schizograpta</i> Müll. Arg.	ME 10%	NS						
<i>Icmadophilaceae</i> Trichel <i>Thamnolia vernicularis</i> (Sw.) Scherter.	ME	2 mg/ml: 67.0%	2 mg/ml: 72.0% 0.2 mg/ml: 36.0%	NVNR	NS			Luo et al. (2006)
<i>Lecanoraceae</i> Körb. <i>Lecanora arata</i> (Hudson) Ach.	AcE, ME, WE	NS	AcE: 94.7% ME: 93.3% WE: 93.2%	AcE: 84.5% ME, WE: NVNR	NS	AcE: 73.0 µg PE ME: 71.0 µg PE WE: 69.8 µg PE (AcE: 54.8 µg RE)	NS	Kosančić and Ranković (2011)

(continued)

Table 1. Continued

Lichen species	Solvent	LP inhibition (IC ₅₀ or %INH)	DPPH (IC ₅₀ or %INH)	SO ⁺ RSA (IC ₅₀ or %INH)	Reducing power (A ₇₀₀ or IC ₅₀)	Phenolic/flavonoid content	Isolated compounds/ composition	Reference
<i>AcE</i>	NS	94.7%	84.5%	ME: 53.7 µg RE WE: 52.6 µg RE	0.109	73.0 µg PE/mg	NS	Ranković et al. (2011)
<i>AcE</i>	NS	52.3%	33.6%	(54.8 µg RE/mg) 43.2 µg PE/mg (34.6 µg RE/mg)	0.061	NS	NS	Ranković et al. (2011)
<i>Lecanora muralis</i> (Schreber) Rabenh.	ME	ME: 87.5% WE: -17.1%	NS	NS	ME: 0.417 WE: 0.233	ME: 87.9 mg GA/g WE: 39.2 mg GA/g	NS	Odabasoglu et al. (2004)
<i>Lobariaceae</i> Chevall. <i>Lobaria pulmonaria</i> (L.) Hoffm.	ME, WE	ME: 87.5% WE: -17.1%	NS	NS	0.098	6.2 mg GA/g	NS	Ranković et al. (2010a)
<i>Nephromataceae</i> Wetm. ex J. C. David & D. Hawksw. <i>Nephroma parile</i> (Ach.) Ach.	ME	ME: 3.6%	NS	NS	0.077	5.6 mg GA/g	NS	Ranković et al. (2010b)
<i>Ochrolechiaceae</i> R. C. Harris ex Lumbsch & I. Schnitt	ME	8.5%	NS	NS	0.202	29.4 mg GA/g	NS	Ranković et al. (2010a)
<i>Ochrolechia parella</i> (L.) A.Massal.	ME	26.6%	NS	NS	ME: 0.215 WE: 0.201	ME: 48.6 mg GA/g WE: 30.3 mg GA/g	NS	Odabasoglu et al. (2005)
<i>Ochrolechia tartarea</i> (L.) A. Massal.	ME	ME: 12.6% WE: 18.4%	NS	NS	NS	NS	NS	Luo et al. (2009)
<i>Parmeliaceae</i> Zenker <i>Bryoria fuscellens</i> (Gyelh.) Brodo & D. Hawksw	ME, WE	ME: >1000 µg/ml ME: >1000 µg/ml	NS	NS	AcE: >1000 µg/ml NVNR	AcE > ME NVNR	NS	Singh et al. (2011)
<i>Cetraria aculeata</i> (Schreber) Fr.	AcE, ME	AcE: 80.3% DmE, EE, ME: NVNR	NS	NS	ME: 0.019 mg/g DmE: 0.048 mg/g EE: 0.009 mg/g	AcE: 0.019 mg/g DmE: 0.048 mg/g EE: 0.009 mg/g	NS	Gülçin et al. (2002)
<i>Cetraria islandica</i> (L.) Ach.	WE	50 µg: 96.0% 100 µg: 99.0% 200 µg: 100.0% 500 µg: 100.0%	NVNR	NVNR	ME: 0.050 mg/g 0.039 µg PE/mg	NS	NS	Kosanić and Ranković (2011)
	AcE, ME, WE	NS	NVNR	NVNR	AcE: 25.0 µg PE ME: 38.0 µg PE WE: 18.2 µg PE	NS	NS	
					(AcE: 7.7 µg RE, ME: 25.5 µg RE, WE: 1.4 µg RE)	32.9 mg GA/g	NS	Ranković et al. (2010b)
<i>Cetraria pinastri</i> (Scop.) Gray	ME	48.8%	NS	NS	0.188	NS	NS	Racine et al. (1980)
<i>Evernia prunastri</i> (L.) Ach.	BE, DE, EE, ME	Crude wax: weak activity DE: activity EE: activity ME: activity	NS	NS	NS	35.5 µg AA/g 0.61 µg Tg	NS	Stojanović et al. (2010)
	ME	NS	727.7 µg/ml	NS	NS	80.7 mg GA/g (27.46 mg rutin/g)	NS	Mitrović et al. (2011)
	ME	NS	>1000 µg/ml	NS	NS	34.1 µg PE/mg	NS	Evernic acid, physodic acid, usnic acid, atranorin, chloroatranorin
	AcE	NS	663.1 µg/ml	1033.6 µg/ml	500 µg/ml: 0.029	NS	NS	Singh et al. (2011)
<i>Flavocetraria nivalis</i> (L.) Kämärfelt & Thell	AcE, DmE, EE, ME	NVNR	NS	NS	AcE: 0.013 mg/g DmE: 0.012 mg/g	NS	NS	Kosanić et al. (2013)

<i>Flavoparmelia caperata</i> (L.) Hale	ME	NS	347.2 µg/ml	NS	21.6 µg AA/g 19.3 µg Tr/g	EE: 0.007 mg/g ME: 0.162 mg/g WE: 11.9 µg GA/mg	NS
	ME	NS	549.0 µg/ml	NS	90.8 mg GA/g (33.6 mg rutin/g)	NS	NS
<i>Hypogymnia physodes</i> (L.) Nyf.	ME	NS	79.7 µg/ml	NS	96.9 µg AA/g 25.3 µg Tr/g	17.5 µg GA/mg	NS
	ME	NS	45.6 µg/ml	NS	141.6 mg GA/g (20.1 mg rutin/g)	AcE: 30.1 µg PE ME: 6.3 µg PE	NS
<i>Letharia lachneriana</i> Krog	Hot WE	NS	AcE: 60.2% ME: 73.2% WE: 30.9%	NS	NVNR ME > AcE > WE	AcE: 20.1 mg PE ME: 6.3 µg PE	Chlororubrocashmeriquinone Rubrocashmeriquinone
<i>Letharia serranderi</i> (Most.) Obermayer	Hot WE	NS	NS	NS	Activity: 163.9	NS	Chlororubrocashmeriquinone Rubrocashmeriquinone 7-Chlorocarotene
<i>Letharia sinensis</i> J.C.Wei & Y.M.Jiang	Hot WE	NS	NS	NS	Activity: 23.8	NS	Rubrocashmeriquinone Chlororubrocashmeriquinone
<i>Parmelia caperata</i> (L.) Ach.	AcE, ME, WE	NS	NVNR AcE > ME > WE	NS	NVNR AcE > ME > WE	AcE: 40.4 µg PE ME: 15.3 µg PE WE: 14.9 µg PE	NS
	AcE	NS	46.4%	61.5%	0.057	40.4 µg PE/mg (15.3 µg RE/mg)	NS
<i>Parmelia centrifuga</i> (L.) Ach.	ME	54.2%	NS	NS	0.375	49.8 mg GA/g	NS
<i>Parmelia cinnita</i> Ach.	ME	16.0%	NS	NS	0.172	12.8 mg GA/g	NS
<i>Parmelia pertusa</i> Schaefer	AcE, ME, WE	NS	NVNR	NVNR Weak activity	Weak activity	AcE: 18.2 µg PE ME: 34.0 µg PE WE: 12.0 µg PE (AcE: 6.6 µg RE, ME: 18.2 µg RE, WE: 5.3 µg RE)	NS
		25.0%	No activity	NS	NS	1.0% (w/w)	NS
<i>Parmelia saxatilis</i> (L.) Ach.	ME	Ferric thiocyanate method WE: 86.1% ME: 94.8% TBA test WE > ME	WE: 19.8% ME: 63.6%	WE: 83.0% ME: 80.0%	ME > WE	WE: 24.3 mg PE/mg ME: 55.1 mg PE/mg	Attranorin, chloroatranorin, salazinic acid
	ME, WE		NS	NS	NS	NS	özen and Kinikoglu (2008)
	AcE	NS	55.3%	35.3%	0.066	40.6 µg PE/mg (20.6 µg RE/mg)	NS
<i>Parmelia sulcata</i> Taylor	ME	NS	493.6 µg/ml	NS	25.3 µg AA/g 41.9 µg Tr/g	7.9 µg GA/g	NS
	ME	NS	584.2 µg/ml	NS	NS	88.3 mg GA/g (44.9 mg rutin/g)	NS
	AcE, ME, WE	NS	NVNR AcE > ME > WE	NVNR AcE > ME > WE	NS	AcE: 38.2 µg PE ME: 25.1 µg PE WE: 9.6 µg PE 18.2 µg PE/mg (6.6 µg RE/mg)	NS
	AcE	NS	38.8%	33.8%	0.034	NS	NS
<i>Parmotrema pseudoincitorum</i> (Abayes) Hale	ME	NS	500 µg/ml: 90.7% + honey: 81.5% 62.1-89.4%	NS	0.265-0.776	Attranorin, lecanoric acid	NS
	BE, AcE	NS	BE: 30.0-65.0% AcE: 35.0-68.0%	NS	NS	Methyl orsellinate, orsellinic acid, atranorin, lecanoric acid	NS
<i>Parmotrema stippeum</i> (Taylor) Hale	ME	NS	NS	NS	NS	NDR	Vema et al. (2008b)

(continued)

Table 1. Continued

Lichen species	Solvent	LP inhibition (IC ₅₀ or %INH)	DPPH (IC ₅₀ or %INH)	SO ⁺ RSA (IC ₅₀ or %INH)	Reducing power (A ₇₀₀ or IC ₅₀)	Phenolic/flavonoid content	Isolated compounds/ composition	Reference	
<i>Parmotrema inlectorum</i> (Delise ex Nyl.) Hale	ME	3.6–71.3% 1.1–57.2% 27.0%	1.1–57.2% 13.6 µg/ml No activity	NS	NS	Depends on lichen culture conditions 1.1 % (W/W)	NS	Güllüce et al. (2006)	
<i>Platismatia glauca</i> (L.) Cath. & C. Cult.	ME, WE	ME: 26.4% WE: 48.2% NS	NS	NS	ME: 0.229 WE: 0.229 NVNR	ME: 75.1 mg GA/g WE: 61.7 mg GA/g AcE: 76.4 mg PE ME: 37.0 µg PE WE: 18.7 µg PE (AcE: 37.6 µg RE, ME: 21.0 µg RE, WE: 1.8 µg RE)	NS	Odabasoglu et al. (2005)	
<i>Pseudovernia furfuracea</i> (L.) Zopf	AcE, ME, WE	AcE: 57.9% WE: 33.9%	AcE: 87.3% ME: 57.9% WE: 33.9%	NS	ME: 37.0 µg PE WE: 18.7 µg PE AcE: 37.6 µg RE, ME: 21.0 µg RE, WE: 1.8 µg RE)	NS	NS	Kosaníć et al. (2011)	
<i>BuE, DemE, EaE, ME</i>	BuE: No activity DemE: 17.8% EaE: 23.8% ME: 17.2% NS	401.7 µg/ml	632.0 µg/ml	500 µg/ml; 0.058	76.4 µg PE/mg	Physodic acid, physodalic acid, atranorin, chloroatranorin, 3-hydroxyphysodic	Attranorin	Givenç et al. (2012)	
<i>Pseudovernia furfuracea</i> (L.) Zopf	AcE	AcE: 82.4% DmE, EE, ME: NVNR	AcE: 51.8% DmE, EE, ME: NVNR	NS	AcE: 0.017 mg/g DmE: 0.012 mg/g EE: 0.001 mg/g ME: 0.027 mg/g	NS	Singh et al. (2013)		
<i>Pseudophlebia pubescens</i> (L.) M. Choisy	AcE, DmE, EE, ME	AcE: NVNR	AcE: >1000 µg/ml ME: >1000 µg/ml AcE: 445.7 µg/ml ME: >1000 µg/ml AcE: 642.6 µg/ml ME: 791.3 µg/ml 667.9 µg/ml	NS	NVNR ME > AcE NVNR AcE = ME NVNR AcE = ME AcE > ME 979.3 µg/ml	ME > AcE NVNR AcE = ME NVNR AcE = ME AcE > ME 31.3 µg PE/mg	NS	Luo et al. (2009)	
<i>Usnea antarctica</i> Du Rietz	AcE, ME	NVNR	AcE: >1000 µg/ml ME: >1000 µg/ml AcE: 445.7 µg/ml ME: >1000 µg/ml AcE: 642.6 µg/ml ME: 791.3 µg/ml 667.9 µg/ml	NS	NVNR AcE > ME NVNR AcE = ME NVNR AcE = ME AcE > ME 979.3 µg/ml	AcE > ME NVNR AcE > ME NVNR AcE = ME AcE > ME 31.3 µg PE/mg	NS	Luo et al. (2009)	
<i>Usnea aurantiacoerulea</i> (Jacq.) Botry	AcE, ME	NVNR	AcE: >1000 µg/ml ME: >1000 µg/ml AcE: 445.7 µg/ml ME: >1000 µg/ml AcE: 642.6 µg/ml ME: 791.3 µg/ml 667.9 µg/ml	NS	NVNR AcE = ME NVNR AcE = ME AcE > ME AcE > ME AcE > ME 979.3 µg/ml	Norstictic acid, usnic acid, atranorin, chloroatranorin Usnic acid, psoromic acid	NS	Luo et al. (2009)	
<i>Usnea barbata</i> Moryka	AcE	NS	WE: 132.4 µg/ml EaE: 125.0 µg/ml ME: 74.6 µg/ml ME: 15.0% WE: 13.4% AcE: 55.0% DmE: 11.0% ME: 87.0% PE: 43.0% 2–20 mg/ml: 3.8–73.3%	NS	WE: 25.0 µg/ml EaE: 25.0 µg/ml ME: 80.0 µg/ml ME: 22.9 µg/ml ME: 80.0 µg/ml NS	ME: 10.5 mg GA/g WE: 10.4 mg GA/g AcE: 14.0 mg/g DmE: 12.0 mg/g ME: 35.0 mg/g PE: 9.0 mg/g 13.0 µg PE/mg	NS	Odabasoglu et al. (2004)	
<i>Usnea complanata</i> (Müller Arg.) Moryka	WE, EE, EaE, ME	WE: 132.4 µg/ml EaE: 157.9 µg/ml ME: 74.6 µg/ml ME: 15.0% WE: 13.4% AcE: 55.0% DmE: 11.0% ME: 73.0% PE: 43.0% NVNR	WE: 25.0 µg/ml EaE: 25.0 µg/ml ME: 80.0 µg/ml ME: 22.9 µg/ml ME: 80.0 µg/ml NS	ME: 0.069 WE: 0.083 NS	ME: 18.0% DmE: 7.0% ME: 56.0% PE: 27.0% 20 mg/ml: 30.0% NVNR	ME: 10.5 mg GA/g WE: 10.4 mg GA/g AcE: 14.0 mg/g DmE: 12.0 mg/g ME: 35.0 mg/g PE: 9.0 mg/g 13.0 µg PE/mg	NS	Behera et al. (2006c)	
<i>Usnea florida</i> (L.) Weber ex F.H.Wigg.	AcE, ME	AcE: 31.0% DmE: 11.0% ME: 73.0% PE: 43.0% 3.8–73.3%	AcE: 18.0% DmE: 7.0% ME: 56.0% PE: 27.0% 20 mg/ml: 30.0%	NS	NS	ME: 10.5 mg GA/g WE: 10.4 mg GA/g AcE: 14.0 mg/g DmE: 12.0 mg/g ME: 35.0 mg/g PE: 9.0 mg/g 13.0 µg PE/mg	NS	Behera et al. (2006c)	
<i>Usnea ghatensis</i> G. Awasthi	AcE, DmE, ME, PE	AcE: 55.0% DmE: 17.0% ME: 87.0% PE: 38.0% 67.0% ME: 82.4% WE: 25.6% 6 mg/ml: 97.3%	AcE: 31.0% DmE: 11.0% ME: 73.0% PE: 43.0% NVNR	AcE: 18.0% DmE: 7.0% ME: 56.0% PE: 27.0% 89.6% NS	NS	ME: 18.0% DmE: 7.0% ME: 56.0% PE: 27.0% 89.6% NS	ME: 10.5 mg GA/g WE: 10.4 mg GA/g AcE: 14.0 mg/g DmE: 12.0 mg/g ME: 35.0 mg/g PE: 9.0 mg/g 13.0 µg PE/mg	NS	Venna et al. (2008a)
<i>Usnea longissima</i> Ach.	ME, WE	ME: 74.0% WE: 26.4%	ME: 74.0% WE: 26.4%	NS	ME: 0.178 WE: 0.100	ME: 38.6 mg GA/g WE: 18.3 mg GA/g 2.62%	NS	Odabasoglu et al. (2004)	
<i>Peltigeraceae</i> Dumort.	ME, WE	ME: 74.0% WE: 26.4%	ME: 74.0% WE: 26.4%	NS	ME: 0.542 WE: 0.218	ME: 66.4 mg GA/g WE: 38.1 mg GA/g	NS	Kim and Cho (2007)	
<i>Peltigera</i> rifescens (Weiss) Humb.	ME	22.6%	NS	NS	NS	27.6 mg GA/g	NS	Odabasoglu et al. (2005)	
<i>Physciaceae</i> Zahbr. <i>Anaptychia ciliaris</i> (L.) Körb.	ME	3.4–52.7%	1.6–53.7%	NS	0.195	27.6 mg GA/g	NS	Rančić et al. (2010a)	
<i>Heterodermia</i> <i>podocarpa</i> (Bel.) D. D. Awasthi	ME	12.7 µg/ml	18.4 µg/ml	NS	Different contents depending on lichen culture conditions	NDR	Venna et al. (2008b)		

<i>Physcia caesia</i> (Hoffm.) Furmr.	AcE, DmE, EE, ME	NVNR	NS	AcE: 0.065 mg/g EE: 0.015 mg/g ME: 0.128 mg/g DmE: 0.016 mg/g	Atranorin, zeorin	Singh et al. (2011)
<i>Ramalinaceae</i> C. Agardh						
<i>Ramalina conduplicans</i> Vain.	ME	NS	58.2–85.4%	NS	NS	Usnic acid, salazinic acid, sekikaic acid
<i>Ramalina hossei</i> Vain.	ME	NS	500 µg/ml: 89.8% + honey: 86.6% No activity	NS	NS	Usnic acid, sekikaic acid
<i>Ramalina pollinaria</i> (Westr.) Ach.	ME	26.0%	No activity	NS	1.0 % (w/w)	Prashith Kekuda et al. (2009)
<i>Ramalina polymorpha</i> (Lilj.) Ach.	ME	19.0%	No activity	NS	0.8 % (w/w)	Güllüce et al. (2006)
<i>Ramalina terebrata</i> Hook. F. & Taylor	ME-WE (90/10)	NS	TLC spray 2–3 antioxidant active spots	NS	NVNR	Bhattarai et al. (2008)
<i>Sphaerophoraceae</i> Fr.						
<i>Sphaerophorus globosus</i> (Huds.) Vain.	AcE, ME	NVNR	AcE: >1000 µg/ml ME: >1000 µg/ml	NVNR AcE = ME	NS	Luo et al. (2009)
<i>Stereocaulaceae</i> Chevall.						
<i>Stereocaulon alpinum</i> Lauer ex Funk	ME-WE (90/10)	NS	TLC spray 2–4 antioxidant active spots	NS	NVNR	Bhattarai et al. (2008)
<i>Teloschistaceae</i> Zahlbr.						
<i>Calopaca regalis</i> (Vain.) Zahlbr.	ME-WE (90/10)	NS	TLC spray 2–3 antioxidant active spots	NS	NVNR	Bhattarai et al. (2008)
<i>Fulgensia fulgens</i> (Sw.) Elenkin	ME	NS	251.7 µg/ml	NS	NS	Paudel et al. (2008)
<i>Xanthoria elegans</i> (Link) Th. Fr.	AcE, DmE, EE, ME	NVNR	NS	0.136	12.4 mg GA/g	Ranković et al. (2010b)
<i>Trypetheliaceae</i> Zenker						
<i>Laurella benguelensis</i> Zahlbr.	ChE and benzene (BF) and methanol fractions (MF)	NS	CHE: 432.0 mg/ml BF: 645.8 mg/ml MF: 758.0 mg/ml	NS	NS	Xhantoria
<i>Umbilicariaceae</i> Chevall.						
<i>Laxatula pastinata</i> (L.) Méat	AcE, ME, WE	NS	AcE: 90.9% ME: 69.9% WE: 65.1%	AcE: 67.4% ME, WE: NVNR WE: 49.6 µg PE WE: 23.9 µg PE	NVNR AcE > ME > WE NS	Manojlović et al. (2010)
<i>Umbilicaria Antartica</i> Frey & Lamb	AcE, ME	NVNR	AcE: 121.3 µg/ml ME: >1000 µg/ml 5 mg/ml: 34.6%	AcE: 91.1% ME: NVNR NS	NVNR AcE > ME NS	Kosanić et al. (2011)
<i>Umbilicaria aprina</i> var. <i>halterifolia</i> Nybl.	ME	NS	60.1%	Moderate 42.0%	Low 0.050	Luo et al. (2009)
<i>Umbilicaria crustulosa</i> AcE	NS	NVNR	NS	39.6 µg PE/mg (28.1 µg RE/mg)	NVNR	Bucukoglu et al. (2012)
<i>(Ach.) Lam</i>						Kosanić et al. (2012b)
<i>Umbilicaria cylindrica</i> (L.) Delise ex Duby	AcE, ME, WE	NS	AcE: 19.1 µg PE ME: 42.3 µg PE WE: 19.1 µg PE	AcE: 42.3 µg PE ME: 42.3 µg PE WE: 19.1 µg PE	NS	Kosanić and Ranković (2011)

(continued)

Table 1. Continued

Lichen species	Solvent	LP inhibition (IC ₅₀ or %INH)	DPPH (IC ₅₀ or %INH)	SO ⁻ RSA (IC ₅₀ or %INH)	Reducing power (A ₇₀₀ or IC ₅₀)	Phenolic/flavonoid content	Isolated compounds/ composition	Reference
<i>Umbilicaria esculenta</i> (Miyoshi) Minks	ME	NS	21.1% (5mg/ml)	NS	NS	(AcE: 12.0 µg RE, ME: 19.1 µg RE, WE: 11.1 µg RE)	NS	Bucukoglu et al. (2012)
<i>Umbilicaria decussata</i> (Vill.) Zahlbr.	AcE, ChE, ME	NS ChE: 29.3µg/ml ME: 35.4µg/ml	39.4% ChE: 31.34µg/ml ME: 34.41µg/ml	Moderate 33.9% NS	Low 0.035 NS	0.9 mg GA/g	19.1 µg PE/mg (12.9 µg RE/mg) ME: 79.2 mg GA/g CHE: 71.3 mg GA/g	Kosanić et al. (2012b); Manojlović et al. (2012b)
<i>Umbilicaria hyperborea</i> (Ach.) Hoffm.	ME	6mg/ml: 92.1%	6 mg/ml: 1.3 mg/ml	NS	NS	1.5%	Salazinic acid, norstictic acid, methyl-β-oryctol carboxylate, ethyl haematomate, usnic acid, atranorin	Kim and Cho (2007)
<i>Umbilicaria leiocarpa</i> DC.	ME	NS	5mg/ml: 23.4%	NS	NS	NS	NS	Bucukoglu et al. (2012)
<i>Umbilicaria nylanderiana</i> (Zahlbr.) H. Magn.	ME	NS	5mg/ml: 32.8%	NS	NS	NS	Gyrophoric acid	Singh et al. (2011)
<i>Umbilicaria polyphylla</i> (L.) Baumg.	AcE	53.0%	400.0 µg/ml	NS	NS	3.0 % (w/w)	AcE: 0.001 mg/g DmfE: 0.029 mg/g EE: 0.001 mg/g ME: 0.046 mg/g 35.9 mg GA/g	Bucukoglu et al. (2012)
<i>Umbilicaria virginis</i> Scheerer	ME	NS	5 mg/ml: 32.9%	NS	NS	NS	NS	Gyrophoric acid
<i>Verrucariaceae</i> Zenker <i>Dermatocarpon inexpectatum</i> (Körber) Hasse	ME, WE	ME: 5.3% WE: 9.5%	NS	ME: 0.119 WE: 0.131	ME: 18.6 mg GA/g WE: 9.2 mg GA/g	NS	Umbilicarinic acid	Bucukoglu et al. (2012)
(Some macrostudies; Macrostudy of 46 lichen species Highest activities: - <i>Scutellinia nylanderiana</i>	ME	S.n.: 85.5% P.p.: 85.4%	S.n.: 90.4% P.p.: 87.8%	NS	S.n.: 1.5 P.p.: 1.0	Sn.: 156.1 µg CE/mg P.p.: 109.3 µg CE/mg	<i>Scutellinia nylanderiana</i> : lecanonic acid	Odabasoglu et al. (2005)
Zahlbr. (S.n.) - <i>Peltigera praetextata</i> (Flörke ex Sommerf.) Zopf (P.p.)				NS	NS	NS	NS	Luo et al. (2010)
Macrostudy of 77 lichen species (<i>Graphidaceae</i> sp.)	DmE, ME	NS	NS	Screening of 77 extracts by two methods. (>50%: <i>Graphina</i> <i>multistriata</i> <i>Graphina</i> <i>salsolinella</i> <i>Graphis</i> <i>assamensis</i> <i>Graphis</i> <i>guimarae</i> <i>Graphis</i> <i>schimperi</i>	NS	NS	NS	Behera et al. (2003)

Macrostudy of 85 lichen species	EE	NS	Screening of 99 extracts. Highest DPPH scavenging activities: <i>Hypogymnia vittata</i> <i>Peltigera aphrodisia</i> <i>Nephromopsis ornata</i> <i>Pseudoevernia furfuracea</i> <i>Cladonia vulcani</i> <i>Peltigera elizobethae</i>	NS	<i>Peltigera aphrodisia</i> : solorinine Hara et al. (2011)
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NS, not studied; NVNR, numeric value not reported; LP, lipoperoxidation; %INH, percentage of inhibition; SO_2^- RSA, superoxide radical scavenging activity; GA, gallic acid; PE, pyrocatechol equivalent; RE, rutin equivalent; AA, ascorbic acid; T, tocopherol; TBA, thiobarbituric acid; CE, catechol equivalent; MDA, malondialdehyde; ChE, chloroform extract; ME, methanol extract; PE, petrol ether extract; AcE, acetone extract; DmE, dimethyl sulfoxide extract; EE, ethanol extract; WE, water extract; DE, diethyl-ether extract; BuE, butanol extract; DcmE, dichloromethane extract; EaE, ethyl acetate extract.

chemical assays, such as DPPH free radical scavenging activity, superoxide anion radical scavenging activity, reducing power, and lipid peroxidation inhibition. Methanol arises as one of the most used solvents for an efficient extraction of lichens bioactive compounds with antioxidant activities and, therefore, many antioxidant activity assays have been performed on methanol extracts (Stojanović et al., 2010; Kosanić & Ranković, 2011; Zambare & Christopher, 2012).

In the last part of our review, we focus on antioxidant responses of these natural products to oxidative stress occurring at intracellular level and *in vivo* trials. Therefore, we collect the more recent investigations on antioxidant activity of lichens species (both purified metabolites and extracts) on *in vitro* cellular substrates (Table 3) and *in vivo* animal models (Table 4).

Apart from all species and assays shown in the tables, some authors have measured similar parameters by other methods (e.g., other radical scavenging properties) and also different parameters related to the antioxidant potential of the aforementioned lichens and compounds. Regarding other radical scavenging properties, Papadopoulou et al. (2007) evaluated the hydroxyl radical scavenging activity of β -orcinol metabolites of the lichen *Hypotrachyna revoluta* (Flörke) Hale (Parmeliaceae); this OH[·]-radical scavenging activity has also been measured in *Toninia candida* (Weber) Th.Fr. (Ramalinaceae) (Manojlovic et al., 2012b), *Usnea ghattensis* G. Awasthi (Parmeliaceae) (Verma et al., 2008a), and *Umbilicaria cylindrica* (L.) Delise ex Duby (Umbilicariaceae) (Manojlovic et al., 2012c). Nitric oxide radical scavenging activity was assayed on *Usnea complanata* (Müll. Arg.), Motyka (Parmeliaceae) (Behera et al., 2012), *Usnea ghattensis* (Verma et al., 2008b), psoromic and usnic acids (Behera et al., 2012), and on other 14 lichen-purified metabolites (Thadhani et al., 2011). Moreover, hydrogen peroxide scavenging activity was investigated on *Parmelia saxatilis* (Özen & Kinalioglu, 2008) and the TEAC value (Trolox equivalent antioxidant capacity) has been determined using the ABTS radical assay in several polar lichen species (Paudel et al., 2008; Singh et al., 2011), as well as in *Usnea ghattensis* (Behera et al., 2005a; Verma et al., 2008a), *Laurera benguelensis* Zahlbr. (Zahlbr.) (Manojlovic et al., 2010), and ramalin compound (Paudel et al., 2011).

Behera et al. (2003, 2006a) assessed the xanthine oxidase inhibitory capacity for many species of the family Graphidaceae; the ferrous ion-chelating activity was investigated in *Umbilicaria cylindrica* (Manojlovic et al., 2012c) and *Toninia candida* (Manojlovic et al., 2012b); and the tyrosinase inhibitory activity has been studied in some lichens and isolated compounds (Behera et al., 2006b; Kim & Cho, 2007; Paudel et al., 2011).

Finally, it is remarkable that the work conducted by Lopes et al. (2008) assayed the radical scavenging properties for many semisynthetic derivatives from the lecanoric acid obtained from a *Parmotrema tinctorum* (Delise ex Nyl.) Hale (Parmeliaceae) specimen and treated with alcohols. They obtained the three most active compounds at scavenging DPPH radical that were orsellinic acid, orcinol and resorcinol, and the lowest activity was displayed by methyl orsellinate.

Table 2. Antioxidant activities of isolated compounds from lichens.

Compound	Origin	LP inhibition (IC ₅₀ or %INH)	DPPH (IC ₅₀ or %INH)	SO RSA (IC ₅₀ or %INH)	Reducing power (A ₇₀₀ or IC ₅₀)	Reference
1,8-Dihydroxy-5-methoxy-3-methylxanthone	<i>Pyrenula japonica</i> Kurok [DE]	Not studied (NS)	Numeric Value Not Reported (NVNR)	NS	NS	Takenaka et al. (2000)
1,5,8-Trihydroxy-3-methylxanthone	<i>Pyrenula japonica</i> Kurok [DE]	NS	NVNR	NS	NS	Takenaka et al. (2000)
1,7-Dihydroxy-3-methylxanthone	<i>Pyrenula japonica</i> Kurok [DE]	NS	NVNR	NS	NS	Takenaka et al. (2000)
1,2,8-Trihydroxy-5-methoxy-3-methylxanthone	<i>Pyrenula japonica</i> Kurok [DE]	NS	NVNR	NS	NS	Takenaka et al. (2000)
1-Chloropanarin	<i>Eriodema chilense</i> Mont.	30%	In rat brain homogen-			Hidalgo et al. (1994)
	at: 27–66%		ate: 7.3–9.5%			
7-Chlorocanarione	<i>Letharia lachneriana</i> Krog [ME]	NS	1/IC ₅₀ = 0.052	Activity: 142.7	NS	Kinoshita et al. (2010)
2-O-Methylsekikaic acid	<i>Ramalina asaphiae</i> W.L. Culb. [ME, AcE]	NS		NS	NS	Valencia-Islas et al. (2007)
Atranol	<i>Letharia l. canariensis</i> (Ach.) Krog [ME]	NYNR		NS	NS	Toledo-Marante et al. (2003)
Atranorin	<i>Placopsis</i> sp.	6.5%	In rat brain homogen-			Hidalgo et al. (1994)
			ate: 7.3–9.5%			
		14.0%	NS	NS	NS	Jayaprakash et al. (2000)
	<i>Parmotrema stippeum</i> (Taylor) Hale [BE, AcE]	Unable to prevent lipid peroxidation	NS	NS	NS	Toledo-Marante et al. (2003)
	<i>Letharia l. canariensis</i> (Ach.) Krog [ME]	NS		NVNR	NS	Lohézic-Le Dévéhat et al. (2007)
	<i>Usnea articulata</i> (L.) Hoffm. [various extracts]					
	<i>Parmotrema stippeum</i> (Taylor) Hale [ME, AcE]	NS	1/IC ₅₀ = 0.576	NS	NS	Valencia-Islas et al. (2007)
	<i>Parmotrema grayana</i> Hue. [DcmE], <i>Heterodermia obscurata</i> (Nyl.) Trevisan [DcmE], and <i>Cladonia</i> sp. [DcmE]	No activity		1.6%	NS	Thadhani et al. (2011)
	<i>Usnea articulata</i> (L.) Hoffm. [various extracts]	NS	NVNR	NVNR	NS	Lohézic-Le Dévéhat et al. (2007)
	<i>Ramalina asaphiae</i> W.L. Culb. & C.F. Culb. [ME, AcE]	1/IC ₅₀ = 0.014	NS	NS	NS	Valencia-Islas et al. (2007)
	<i>Letharia l. serranderi</i> (Motyka) Obermayer [AcE]	NS		Activity: 137.2	NS	Kinoshita et al. (2010)
	<i>Letharia l. canariensis</i> (Ach.) Krog [ME]	50.0–60.0%	NS	NS	NS	Toledo-Marante et al. (2003)
	<i>Letharia l. canariensis</i> (Ach.) Krog [ME]	50.0–57.0%	NS	NS	NS	Toledo-Marante et al. (2003)
	<i>Parmotrema stippeum</i> (Taylor) Hale [ME, AcE]	NS	1/IC ₅₀ = 0.589	NS	NS	Valencia-Islas et al. (2007)
	<i>Letharia l. canariensis</i> (Ach.) Krog [ME]	NYNR	NS	NS	NS	Toledo-Marante et al. (2003)
	<i>Letharia l. serranderi</i> (Motyka) Obermayer [AcE], <i>L. sinensis</i> J.C.Wei & Y.M.Jiang [ME], and <i>L. cashmeriana</i> Krog [ME]	NS		Activity: 277.4	NS	Kinoshita et al. (2010)
	<i>Usnea articulata</i> (L.) Hoffm. [various extracts]	NS	18.0%	NVNR	NS	Lohézic-Le Dévéhat et al. (2007)
	<i>Usnea longissima</i> Ach. [EE]	NYNR – no antioxidant activity	NS	NS	NS	Ataley et al. (2011)
	<i>Protousnea magellanica</i> (Mont.) Krog	NS	No scavenging activity	NS	NS	Brisideli et al. (2013)
	<i>Protousnea malacea</i> (Srift) Krog	8.6%	NS	NS	NS	Hidalgo et al. (1994)
		In rat brain homogen-				
		ate: 1.7–8.4%				
Cryptostictinolide			27.0%	92.5%	NS	Thadhani et al. (2011)
Diffractaic acid			NVNR	NDR	NS	Lohézic-Le Dévéhat et al. (2007)
Divaricatic acid						
Ergosterol peroxide						
Erythrin	<i>Lobaria pulmonaria</i> L. (Hoffm.) [AcE]	LNVNR	NVNR	NS	NS	Ataley et al. (2011)
Ethyl chlorohematommate	<i>Roccella montagnei</i> Bel. [AcE]	NS	No scavenging activity	84.1%	NS	Thadhani et al. (2011)
Ethyl hematommate	<i>Letharia l. canariensis</i> (Ach.) Krog [ME]	50.0–60.0%	NS	NS	NS	Toledo-Marante et al. (2003)
Evemic acid	<i>Evernia prunastri</i> Ach. [AcE]	NVNR	NS	NS	NS	Toledo-Marante et al. (2003)
Fumaroproctoeric acid	<i>Usnea articulata</i> (L.) Hoffm. [various extracts]	NS	322.4 µg/ml	561.8 µg/ml	NS	Kosanić et al. (2013)
			20.0%	566.0 µM	NS	Lohézic-Le Dévéhat et al. (2007)

Glyphoric acid		Büyükoglu et al. (2012)
Hematomic acid		Toledo-Marante et al. (2003)
Isidiphorin		Atabay et al. (2011)
Lecanonic acid		Jayaprakash et al. (2000)
<i>Umbilicaria virginis</i> Schaefer	NS	NS
<i>Letharia canariensis</i> (Ach.) Krog [ME]	50.0-60.0%	NS
<i>Lobaria pulmonaria</i> L. (Hoffm.) [AcE]	NVNR	NVNR – high scavenging activity
<i>Parmotrema stippeum</i> (Taylor) Hale [BE, AcE]	12.0-36.0%	NS
<i>Parmotrema tinctorium</i> (Nyl.) Hale	NS	50.9%
<i>Sictia nylanderiana</i> Zahlbr. [ME]	51.5%	NS
<i>Parmotrema grayana</i> Hue. [ME]	NS	98.4%
<i>Umbilicaria apirina</i> var. <i>halei</i> Nyl. [ME]	34.0%	NS
<i>Cladonia</i> sp. [ME]	NS	32.5%
<i>Stereocaulon applanatum</i> Laur.	NS	No scavenging activity
<i>Letharia canariensis</i> (Ach.) Krog [ME]	NS	No scavenging activity
<i>Letharia canariensis</i> (Ach.) Krog [ME]	66.0-70.0%	NS
<i>Parmotrema grayana</i> Hue. [DcmE] and <i>Heterodermia obscurata</i> (Nyl.) Trevisan [DcmE]	NVNR	NS
<i>Usnea articulata</i> (L.) Hoffm. [various extracts]	NS	NVNR
<i>Heterodermia obscurata</i> (Nyl.) Trevisan [DcmE], and <i>Claudonia</i> sp. [DcmE]	NS	No scavenging activity
<i>Parmotrema stippeum</i> (Taylor) Hale [BE, AcE]	18.0-40.0%	NS
<i>Letharia canariensis</i> (Ach.) Krog [ME]	NVNR	1.5%
<i>Parmotrema grayana</i> Hue. [ME], <i>Heterodermia obscurata</i> (Nyl.) Trevisan [DcmE], and <i>Claudonia</i> sp. [ME]	NS	NS
<i>Roccella montagnei</i> Bell. [AcE]	NS	NS
<i>Usnea articulata</i> (L.) Hoffm. [various extracts]	NS	NS
<i>Toninia candida</i> (Weber Th. Fr. [AcE]	NS	NS
<i>Parmotrema grayana</i> Hue. [ME]	NS	NS
<i>Parmotrema stippeum</i> (Taylor) Hale. [BE, AcE]	26-50%	NS
<i>Parmotrema grayana</i> Hue. [ME]	NS	133.5 µg/ml
<i>Erioderma ciliense</i> Mont.	23.0%	49.7%
In rat brain homogenate: 13.0-36.0%	NS	38.2%
<i>Pseudoevernia furfuracea</i> (L.) Zopf. [AcE]	NS	NS
<i>Cornicularia aculeata</i> (Schreb.) Ach.	NS	500 µg/ml: 0.547
<i>Parmelia caperata</i> (L.) Ach. [AcE]	NS	NS
<i>Usnea complanata</i> (Müll. Arg.) Motschka [total extract]	0.2 mg/ml	NS
<i>Lobaria pulmonaria</i> L. (Hoffm.) Ach. f. & Taylor. [ME]	NVNR	7.9%
<i>Ramalina terebra</i> Hook. f. & Taylor. [ME]	NS	NS
Physodic acid	69.1 µg/ml	500 µg/ml: 0.664
Protolichesterinic acid	0.990 µg/ml	0.4 µg of BHT equivalent
Protrrocetrinic acid	No scavenging activity	NS
Psoromic acid	138.2 µg/ml	NS
Pulmonaritin	0.271 µg/ml	500 µg/ml: 0.090
Ramalin	NS	NS
Rhizonaldehyde	118.2 µg/ml	NS
Rhizonyl alcohol	10.2 µg/ml	NS
Rubrocashmerquinone	177.6 µg/ml	NS
Salazinic acid	0.271 µg/ml	NS
<i>Heterodermia obscurata</i> (Nyl.) Trevis. [ME]	NVNR	Activity: 87.2
<i>Peltigera aphlophora</i> (L.) Willd. [EE]	NVNR	NS
<i>Usnea articulata</i> (L.) Hoffm. [various extracts]	NVNR	NS
<i>Lobaria pulmonaria</i> L. (Hoffm.) [AcE]	NVNR	Kinoshita et al. (2010)
<i>L. seminervosa</i> (Motschka) Obermayer [AcE], and <i>L. sinensis</i> Wei & Jiang. [ME]	NVNR	Manojlovic et al. (2012a)
<i>Parmelia</i> sp. (<i>P. saxatilis</i> (L.) Ach. and <i>P. sulcata</i> Taylor.) [AcE]	NS	Atabay et al. (2011)
<i>Parmotrema stippeum</i> (Taylor) Hale, [ME, AcE]	NS	Valencia-Islas et al. (2007)
<i>Heterodermia obscurata</i> (Nyl.) Trevis. [ME]	NS	Thadhani et al. (2011)
<i>Peltigera aphlophora</i> (L.) Willd. [EE]	NS	Hara et al. (2011)
<i>Usnea articulata</i> (L.) Hoffm. [various extracts]	NVNR	Lohézic-Le Dévéhat et al. (2012)
<i>Lobaria pulmonaria</i> L. (Hoffm.) [AcE]	NVNR	Atabay et al. (2011)
<i>Umbilicaria nylanderioides</i> (Zahlbr.) H. Magn [ME]	NS	Büyükoglu et al. (2012)
Sekikaic acid	32.6%	NS
Solorinine	120.0 µmol/ml	NS
Stictic acid	NVNR	NS
Umbilicanic acid	68.1%	NS

Table 2. Continued

Compound	Origin	LP inhibition (IC ₅₀ or %INH)	DPPH (IC ₅₀ or %INH)	SO RSA (IC ₅₀ or %INH)	Reducing power (A ₇₀₀ or IC ₅₀)	Reference
Usnic acid	<i>Letharia canariensis</i> (Ach.) Krog [ME]	Unable to prevent lipid peroxidation	NS	NS	NS	Toledo-Marante et al. (2003)
	<i>Usnea articulata</i> (L.) Hoffm.[various extracts]	NS	NVNR	NVNR	NS	Lohézic-Le Dévéhat et al. (2007)
	<i>Ramalina ascalinae</i> W. L. Culb. & C. F. Culb. [ME, AcE]	NS	1/IC ₅₀ = 0.112	NS	NS	Valencia-Islas et al. (2007)
	<i>Usnea longissima</i> Ach. [EE]	NVNR – no antioxidant activity	No scavenging activity	NS	NS	Atalay et al. (2011)
	<i>Parmotrema grayana</i> Hue. [DcmE]	NS	No scavenging activity	59.9%	NS	Thadhani et al. (2011)
	<i>Usnea corollinata</i> (Müll. Arg.) Molyka [Total extract]	0.214 mg/ml	0.195 mg/ml	NS	NS	Behera et al. (2012)
	<i>Parmelia caperata</i> (L.) Ach. [AcE]	NS	60.7 µg/ml	97.3 µg/ml	500 µg/ml: 0.547	Manojlović et al. (2012a)
	<i>Usnea barbata</i> Molyka [AcE]	NS	130.7 µg/ml	197.3 µg/ml	500 µg/ml: 0.664	Rankovic et al. (2012)
Variolanic acid	<i>Cladonia lepidophora</i> Alti & Kashiw [Ochrolecia deceptoria] (Hue) Darb.	NS	No scavenging activity	NS	NS	Brisdelli et al. (2013)
	<i>Lobaria pulmonaria</i> L. (Hoffm.) [AcE]	NS	No scavenging activity	NS	NS	Atalay et al. (2011)
Vesuvianic acid	<i>Psoroma pallidum</i> Nyg [DcmE]	NVNR	NS	NS	NS	Brisdelli et al. (2013)
Vicanicin	<i>Cladonia</i> sp. [DcmE]	NS	No scavenging activity	1.6%	NS	Thadhani et al. (2011)

NS, not studied; NVNR, numeric value not reported; LP, lipoperoxidation; %INH, percentage of inhibition; SO RSA, superoxide radical scavenging activity; GA, gallic acid; PE, pyrocatechol equivalent; RE, rutin equivalent; AA, ascorbic acid; T, tocopherol; TBA, thiobarbituric acid; CE, catechol equivalent; MDA, malodialdehyde; CHE, chloroform extract; ME, methanol extract; PE, petrol ether extract; ACE, acetone extract; DmE, dimethyl sulfoxide extract; EE, ethanol extract; BE, benzene extract; DE, diethyl-ether extract; WE, water extract; BuE, butanol extract; DcmE, dichloromethane extract; EaE, ethyl acetate extract.

Table 3. Evaluations of antioxidant parameters on cellular substrates.

Lichen specie/isolated compound	Solvent/origin	Cell line	LP inhibition	Antioxidant activity markers	Reference
Atranorin	Comercial	Human neuron-like cells (SH-SY5Y)	↑ Peroxyl radical-induced lipoperoxidation <i>in vitro</i>	Good antioxidant capacity in TRAP/TAR assays ↑ H ₂ O ₂ and NO production and SO [·] - scavenging activity Protection of SH-SY5Y cells against H ₂ O ₂ -induced cell viability impairment	Melo et al. (2011)
<i>Cetraria islandica</i> (L.) Ach. Diffractaic acid	Methanol <i>Protosyne magellanica</i> (Mont.) Krog	Human lymphocytes HeLa cell line	↓ MDA levels Not studied (NS)	↑ SOD and GPx activities No effect on intracellular ROS level	Kotan et al. (2011)
<i>Evernia prunastri</i> (L.) Ach. Lobaric acid	Methanol <i>Stereocaulon alpinum</i> Lauter ex Funk	Human lymphocytes HeLa cell line	↓ MDA levels NS	No protection against t-BHP-induced increase in intracellular ROS level ↑ GSH levels, SOD and GPx activities No effect on intracellular ROS level No protection against t-BHP-induced increase in intracellular ROS level	Brisdelli et al. (2013)
<i>Peltigera horizontalis</i> (Hudson) Baumg. <i>Peltigera praetextata</i> (Flörke ex Sommerf.) Zopf	Methanol	Human lymphocytes	↓ MDA levels	↑ GSH levels, SOD and GPx activities ↑ GSH levels, SOD and GPx activities	Alpsoy et al. (2015) Brisdelli et al. (2013)
	Methanol	Human lymphocytes	↓ MDA levels	↑ GSH levels, SOD and GPx activities	Nardemir et al. (2013)
	Methanol	Human lymphocytes	↓ MDA levels	↑ GSH levels, SOD and GPx activities	Nardemir et al. (2013)

Protolichesterinic acid	<i>Corniculularia aculeata</i> (Schreb.) Ach.	HeLa cell line	NS	Brisdelli et al. (2013)
Ramalin	<i>Ramalina terebrata</i> Hook. f. & Taylor. [ME]	Murine macrophage Raw 264.7 cells	NS	Paudel et al. (2011)
Salazinic acid	<i>Xanthoparmelia cantschadalis</i> (Ach.) Hale [ME]	Astrocyte cell line U373-MG	NS	de Paz et al. (2010)
Stictic acid	<i>Xanthoparmelia conspersa</i> (Ach.) Hale [ME]	Astrocyte cell line U373-MG	NS	de Paz et al. (2010)
<i>Umbilicaria vellea</i> (L.) Ach.	Methanol	Human lymphocytes	↓ MDA levels	Aslan et al. (2011)
Usnic acid (UA)	<i>Xanthoparmelia conspersa</i> (Ach.) Hale and <i>X. cantschadalis</i> (Ach.) Hale [ME]	Astrocyte cell line U373 MG	NS	de Paz et al. (2010)
Variolaric acid	Comercial	Human neuron-like cells (SH-SY5Y)	↑ Lipoperoxidation	Rabelo et al. (2012)
Usnic acid (UA)	<i>Cladonia lepidophora</i> Ahti & Kashiw.	HeLa cell line	NS	Brisdelli et al. (2013)
Vicanicin	<i>Ochrolechia decepcionis</i> (Hue) Darb.	HeLa cell line	NS	Brisdelli et al. (2013)
<i>Xantho somloensis</i> (Gyelnik) Hale.	Methanol	Human lymphocytes	↓ MDA levels	No protection against t-BHP-induced increase in intracellular ROS level
<i>Xanthoparmelia cantschadalis</i> (Ach.) Hale	Methanol	Astrocyte cell line U373 MG	NS	No protection against t-BHP-induced increase in intracellular ROS level
<i>Xanthoparmelia conspersa</i> (Ach.) Hale	Methanol	Astrocyte cell line U373 MG	NS	No protection against t-BHP-induced increase in intracellular ROS level
<i>Xanthoria elegans</i> (Link) Th. Fr.	Water	Human lymphocytes	NS	Turkez et al. (2011)

TBARS, thiobarbituric acid reactive substances; SOD, superoxide dismutase; GPx, glutathione peroxidase; GSH, reduced glutathione; AFB₁, aflatoxin B₁; AE, water extract; ME, methanol extract; ROS, reactive oxygen species; OH, hydroxyl radical; CAT, catalase; TAC, total antioxidant capacity; TOS, total oxidative stress; MPx, myeloperoxidase; cNOS, constitutive nitric oxide synthase; iNOS, inducible nitric oxide synthase; GR, glutathione reductase; GST, glutathione transferase; DE, diethyl ether extract; AcE, acetone extract.

Table 4. *In vivo* evaluations of antioxidant activities.

Lichen species/isolated compound	Solvent/origin	LP inhibition	Antioxidant enzymes and other antioxidant markers	Reference
<i>Cetraria islandica</i> (L.) Ach.	—	↓ MDA levels	↑ GSH levels ↓ CAT and GPx activities Association of magnesium augmented the antioxidant effect	Cernescu et al. (2011)
Diffractaic acid	<i>Usnea longissima</i> Ach. [DE]	↓ lipoperoxidation level in tissues	↑ SOD and GPx activities and GSH levels ↓ CAT and MPx activities	Bayir et al. (2006)
Fumarprotocetratic acid	<i>Cladonia verticillaris</i> (Raddi) Fr. [AcE] Methanol	↓ endotoxin-induced lipid peroxidation ↓ lipoperoxidation level in tissues	↑ cNOS activity and ↓ iNOS activity Not studied (NS)	de Barros Alves et al. (2014)
<i>Lobaria pulmonaria</i> (L.) Hoffm.	Methanol	↓ lipoperoxidation	↑ SOD, GPx and GSH levels in tissues No effect on CAT and MPx levels	Karakus et al. (2009)
<i>Peltigera rufescens</i> (Weiss) Humb.	Methanol	↓ lipoperoxidation	↑ SOD, CAT, GSH, GR and GPx levels	Tanas et al. (2010)
<i>Usnea ghattensis</i> G. Awasthi	Methanol	↓ MDA formation in liver tissue	↓ MPx and iNOS activities	Varma et al. (2008a)
<i>Usnea longissima</i> Ach.	Water	47.1% inhibition	Depletion in antioxidant enzymes (SOD, CAT, GPx) and GSH ↑ SOD and GST levels ↓ CAT activity.	Halici et al. (2005)
Usnic acid	<i>Usnea longissima</i> Ach. [DE]	↓ lipoperoxidation level in tissues	Reducing power (A ₇₀₀): 0.1; phenolic content: 18.3 mg GA/g ↑ SOD, GSH and GPx levels ↓ CAT, GR and MPx activities ↑ cNOS activity and ↓ iNOS activity	Odabasoglu et al. (2006)

TBARS, thiobarbituric acid reactive substances; SOD, superoxide dismutase; GPx, glutathione peroxidase; GSH, reduced glutathione; CAT, catalase; TAC, total antioxidant capacity; AFB₁, aflatoxin B₁; AE, water extract; ME, methanol extract; ROS, reactive oxygen species; OH, hydroxyl radical; MPx, myeloperoxidase; TOS, total oxidative stress; MPx, total oxidative stress; MPx, myeloperoxidase; cNOS, constitutive nitric oxide synthase; iNOS, inducible nitric oxide synthase; GR, glutathione reductase; GST, glutathione transferase; DE, diethyl ether extract; AcE, acetone extract.

Conclusions

The present review reports the biological activity of more than 75 different lichen species, as well as more than 65 purified metabolites, isolated from these or other species. The study of their antioxidant activities has recently been started and they have been determined by various chemical *in vitro* assays as first approach, with some of them showing interesting results. Further knowledge of this potential implies deeper research on their activities in order to understand the implied mechanisms. Thus, in this report, we also reflect the few available data about *in vitro* antioxidant activities of some lichen species and purified metabolites on cellular substrates and *in vivo* on animal models.

Concerning antioxidation, the most interesting compounds are polyphenols. The antioxidant properties of polyphenols are due to the presence of their many phenolic hydroxyl groups, which confer high potential for scavenging free radicals (Dai & Mumper, 2010; Sawa et al., 1999). For instance, phenolic compounds are able to donate hydrogen to reactive radicals and break the chain reaction of lipid oxidation at the initiation step (Gülçin et al., 2004).

Then, the strong antioxidant activity shown by some lichen extracts or metabolites, and assessed by different systems, can be attributed to their high total polyphenolic contents (specially depsides, depsidones, dibenzofurans, etc.), since a positive correlation between phenolic composition and antioxidant activity has been proved for most of them (Kosanić et al., 2011; Manojlović et al. 2012c); at least, it suggests that polyphenols might be the major antioxidant compounds in studied lichens. Nevertheless, there have been other studies in which results did not show any positive correlation between antioxidant activity of certain lichens and total phenolic contents (Odabasoglu et al., 2004; Stojanović et al., 2010). This fact implies that other minor compounds should not be ignored but antioxidant activity might be as well attributed to the presence of non-phenolic compounds, antagonistic or synergistic interactions between constituents, and even distinct antioxidant activities of individual phenolics.

In the previous tables, the great diversity of lichens and their substances is shown, and one might deduce that the increasing interest in the study of its pharmacological properties is promoting further phylogenetic studies in an evolutionary context. Based on molecular data mainly, they are leading to a more complex classification of lichen families and species (Crespo et al., 2010). Moreover, phylogenetic analysis of biosynthetic genes can facilitate the discovery of novel compounds, novel genes, and, therefore, unknown producers of pharmaceutical relevant compounds, including antioxidants: the greatest challenges would be to find the biosynthetic gene of interest or assign function to each of the biosynthetic genes found in a lichen genome (Schmitt & Barker, 2009). Considering the difficulties still found for the *in vitro* culture of lichens and different culture conditions result in different antioxidant activities of lichen extracts (due to the production of different amount and type of secondary metabolites depending on culture characteristics) (Behera et al., 2005b), a global approach to the lichen metabolomic features seems to be crucial for the development of new and viable biotechnological processes. These will allow

production of suitable amounts of unique isolated antioxidant compounds from lichens (Boustie & Grube, 2005).

Through this review of literature, we can conclude that lichens are a potential source of natural antioxidants but, at the same time, there is still a need for a deeper research in order to establish their possibilities and a better understanding of their mechanisms of action. This goal can be achieved by the better isolation of purified metabolites and more studies on appropriate cell lines and *in vivo*, in order to identify molecular targets, active compounds, and structure–activity correlations.

Declaration of interest

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