MINI-REVIEW

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Medicinal mushrooms as a source of antitumor and immunomodulating polysaccharides

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Abstract The number of mushrooms on Earth is esti- various allogeneic and syngeneic tumors, and prevent tutantly for modern medicine, they represent an unlimitedintact T-cell component; their activity is mediated mushrooms contain biologically active polysaccharides erties, but also on biotechnological availability. The in fruit bodies, cultured mycelium, culture broth. Data on present review analyzes the pecularities of polysacchamushroom polysaccharides have been collected from rides derived from fruiting bodies and cultured mycelihigher Hetero- and Homobasidiomycetes. These poly- tion today) in selected examples of medicinal mushsaccharides are of different chemical composition, with rooms. most belonging to the group oβ-glucans; these have β -(1 3) linkages in the main chain of the glucan and additional β -(1 6) branch points that are needed for Introduction their antitumor action. High molecular weight glucans weight. Chemical modification is often carried out to im- kind as an edible and medical resource. A number of biocarboxymethylation. Most of the clinical evidence for ing properties (Mizuno 1996, 1999a, b, 2002; Lorenzen antitumor activity comes from the commercial polysac- and Anke 1998; Borchers et al. 1999; Ooi and Liu 1999; charides lentinan, PSK (krestin), and schizophyllan, but Wasser and Weis 1999; Tzianabos 2000; Reshetnikov polysaccharides of some other promising medicinal et al. 2001). Historically, hot-water-soluble fractions (demushroom species also show good results. Their activity coctions and essences) from medicinal mushrooms, i.e.,

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mated at 140,000, yet maybe only 10% (approximately mor metastasis. Polysaccharides from mushrooms do not 14,000 named species) are known. Mushrooms comprise attack cancer cells directly, but produce their antitumor a vast and yet largely untapped source of powerful new effects by activating different immune responses in the pharmaceutical products. In particular, and most impor- host. The antitumor action of polysaccharides requires an source of polysaccharides with antitumor and immuno-through a thymus-dependent immune mechanism. Practistimulating properties. Many, if not all, Basidiomycetes cal application is dependent not only on biological prop-651 species and 7 infraspecific taxa from 182 genera of um (the two main methods of biotechnological produc-

appear to be more effective than those of low molecular For millennia, mushrooms have been valued by humanprove the antitumor activity of polysaccharides and their active molecules, including antitumor substances, have clinical qualities (mostly water solubility). The main been identified in many mushroom species. Polysacchaprocedures used for chemical improvement are: Smith rides are the best known and most potent mushroomdegradation (oxydo-reducto-hydrolysis), formolysis, and derived substances with antitumor and immunomodulatis especially beneficial in clinics when used in conjunc- mostly polysaccharides, were used as medicine in the Far tion with chemotherapy. Mushroom polysaccharides pre-East, where knowledge and practice of mushroom use vent oncogenesis, show direct antitumor activity against primarily originated (Hobbs 1995, 2000). Mushrooms

such as Ganoderma lucidum (Reishi), Lentinus edodes (Shiitake), Inonotus obliquus (Chaga) and many others have been collected and used for hundreds of years in Korea, China, Japan, and eastern Russia. Those practices still form the basis of modern scientific studies of fungal medical activities, especially in the field of stomach, prostate, and lung cancers. It is notable and remarkable how reliable the facts collected by traditional eastern medicine are in the study of medicinal mushrooms (Ying et al.

1987; Hobbs 1995, 2000; Wasser and Weis 1997a, b, 1999; Stamets 2000).

Ikekawa et al. (1969) published one of the first scientific reports on antitumor activities of essences obtained nition of Chang and Miles (1992): 'a macrofungus with a mals (Ikekawa et al. 1982, 1992; Ikekawa 2001). Soon from the Dictionary of the Fungi is at least 14,000 and thereafter the first three major drugs were developed fromperhaps as many as 22,000 (Hawksworth 2001). However, biomass of Trametes versicolor (Turkwey Tail), lentinan the liquid cultured broth product oschizophyllum commune(Split Gill). For almost 40 years, medicinal mushrooms have been intensively investigated for medicinal effects in in vivo and in vitro model systems, and many pics, especially those species forming ectomycorrhizas new antitumor and immunomodulating polysaccharides with native trees. In various tropical areas, 22–55% (in have been identified and put into practical use (Mizuno some cases up to 73%) of mushroom species have 1996, 1999a; Wasser and Weis 1999; Ikekawa 2001).

among higher Basidiomycetes mushrooms, and most ofhave been described and catalogued in thendex of them have unique structures in different species. More-Fungi in the 10 years from 1990 to 1999 revealed that over, different strains of one Basidiomycetes species can about 60% of all newly described fungi are from the troproduce polysaccharides with different properties. For ex-pics (Hawksworth 1993, 2001), and this is also the case from the strain Trametes (Coriolus) versicolor CM-101, covered in Europe and North America. whereas polysaccharide-peptide (PSP) in China was decharide component but with different protein moleculexies', i.e., populations functioning as separate biological bound to the polysaccharide (Hiroshi and Takeda 1993). species but covered by a single scientific name. A single

ing polysaccharides from higher Basidiomycetes mush- more biological species (Hawksworth 2001). rooms are analyzed. More attention is given to their common features than to specific pecularities. The re- number of fungi on Earth range from 500,000 to 9.9 milview summarizes the general state of knowledge in the lion species, of which only 80,060 are named. A working area of biodiversity of mushrooms and their polysaccha-figure of 1.5 million species is generally accepted, and ble ways of chemical modification; results of experimen- of which maybe only 10% are known. Meanwhile, of ture mycelia in selected examples of studied medicinal (Chang 1999; Wasser and Weis 1999; Reshetnikov et al. mushrooms.

The vast quantity and diversity of mushrooms with antitumor polysaccharides

The total number of described fungi of all kinds is curbeen investigated for biologically active polysaccharently at least 80,060 species; a figure based on the total rides, and most of them possess such substances. At derived from addition of the numbers of species in each least 651 species and 7 infraspecific taxa representing Fungi (Kirk et al. 2001). This figure includes all organ-

chromistan fungi, chytridiaceous fungi, lichen-forming fungi, filamentous fungi, molds, and yeasts.

By the term 'mushrooms', we generally mean the defifrom fruiting bodies of mushrooms belonging to the fami- distinctive fruiting body which can be either hypogeous or ly Polyporaceae (Aphyllophoromycetideae) and a few epigeous, large enough to be seen with the naked eye and other families, manifested as host-mediated activity to be picked by hand'. The number of filamentous fungi against grafted cancer - such as Sarcoma 180 - in ani- that are mushrooms in the sense of this definition deduced medicinal mushrooms. All three were polysaccharides, the real number of such species on Earth is undoubtedly specifically β-glucans: krestin from cultured mycelial much higher. Two main reasons for the real number being higher are (1) the great number of as yet undescribed spefrom fruiting bodies of L. edodes, and schizophyllan from cies and (2) the fact that many morphologically defined mushroom 'species' prove to be assemblages of several biological species (Hawksworth 2001).

Most new mushrooms are being discovered in the troproved to be undescribed (Hawksworth 2001). An analy-Biologically active polysaccharides are widespread sis of the localities from which fungi new to science ample, the proteoglucan krestin was developed in Japanfor mushrooms, although new species continue to be dis-

Studies of compatability and molecular sequences beveloped in submerged culture of the strain Cov-1 of the tween mushrooms previously considered to be the same same species. Both proteoglucans have the same polysac-species on morphological grounds revealed 'cryptic spe-In the present review, antitumor and immunomodulat-morphologically defined species may consist of 20 or

Taking all this into account, recent estimates of the rides; the chemical structure of polysaccharides and its new data suggests that this is not unreasonable. The connection with their antitumor activity, including possi- number of mushrooms on Earth is estimated at 140,000, tal testing and clinical use of antitumor or immunostimu- those ~14,000 species that we know today, about 50% lating polysaccharides; possible mechanisms of their bio- are considered to possess varying degrees of edibility, logical action; and, finally, the difference in polysaccha- more than 2,000 are safe, and about 700 species are ride fraction composition in fruiting bodies and pure cul- known to possess significant pharmacological properties 2001). Thus, it is clear that mushrooms represent a major and as yet largely untapped source of powerful new pharmaceutical products.

Higher Basidiomycetes mushrooms represent an unlimited source of antitumor or immunostimulating polysaccharides. All main taxonomic mushroom groups have genus given in the latest edition of the ictionary of the 182 genera of Hetero- and Homobasidiomycetes mushrooms contain antitumor or immunostimulating polyisms traditionally studied by mycologists: slime molds, saccharides, as is evident from Table 1 (adapted from

Table 1 Higher Basidiomycetes mushrooms containing antitumor or immunostimulating polysaccharides

Taxa (number of species studied)	A ctivity ag	gainst:	Source	
	Sarcoma 180 solid cancer	Ehrlich solid cancer		
Heterobasidio mycetes				
Auriculariales – Auricularia (3)	70–90	60–80	Ohtsuka et al. 1973; Ukai et al. 1982; Song et al. 1998	
Dacrymycetales – Calocera (1) Dacrymyces (1) Tremellales – Exidia (1) Guepinia (1) Holtermannia (1) Phlogiotis (1) Protodaedalea (1) Pseudohydnum(1) Tremella (2) Tremellodon (1)	60–90 60–100	60 70–100	Ohtsuka et al. 1973 Ohtsuka et al. 1973; Gao et al. 1997	
Homobasidiomycetes				
Aphyllophoromycetideae				
Cantharellaceae – Cantharellus (5) Craterellus (2) Clavariaceae – Clavaria (4) Clavariadeiphus (2) Clavulinopsis (4) Lentaria (1) Clavulinaceae – Clavulina (1) Sparassidaceae – Sparassis (1)	60–100 60–90 70–90 100	60–90 60–100 80 100	Ohtsuka et al. 1973 Ohtsuka et al. 1973 Ohtsuka et al. 1973 Ohtsuka et al. 1973; Ohno et al. 2000; Yadomae and Ohno 2000	
Ramariaceae – Ramaria (5)	60-80	60–70	Ohtsuka et al. 1973	
Hydnaceae – Hydnum(1)	70	90	Ohtsuka et al. 1973; Chung et al. 1982	
Hericiaceae – Echinodontium(2) Hericium (2) Laxitextum(1)	70–90	60–80	Ohtsuka et al. 1973; Mizuno 1999b	
Corticiaceae – Aleurodiscus (1) Cotylidia (2) Laxitextum (1) Lopharia (1) Merulius (2) Phlebia (2) Sarcodontia (1) Sistotrema (1) Steccherinum (1) Stereum (13)	60–100	60–100	Ohtsuka et al. 1973	
Coniophoraceae –Serpula (1)	70	60	Ohtsuka et al. 1973	
ThelephoraceaeBankera (1) Calodon (4) Hydnellum(2) Polyozellus (1) Sarcodon (2) Thelephora (1)	60–100	70–100	Ohtsuka et al. 1973; Song et al. 1998; Mizuno 2000	
Hymenochaetaceae -Coltricia (4) Cryptoderma(6) Cyclomyces(1) Fuscoporia (1) Hymenochaete(4) Hymenostilbe(1) Inonotus(6) Onnia (1) Phellinus (6) Pyrrhoderma (1)	60–100	90–100	Ohtsuka et al. 1973; K im et al. 1996; Han et al. 1999; Mizuno 2000	
Fistulinaceae – Fistulina (2)	80	90	Ohtsuka et al. 1973; Ueno et al. 1978	
Ganodermataceae -Ganoderma (7)	70–100	70–100	Ohtsuka et al. 1973; Nakashima et al. 1979 Miyazaki and Nishijima 1981; Ukai et al. 1983; Zhang and Lin 1999	
Polyporaceae – Amauroderma(1) Coriolellus (1) Coriolus (8) Cymatoderma(2) Cystidiophorus (1) Daedalea (1) Daedaleopsis (3) Dendropolyporus (1) Favolus (3) Fomes (2) Fomitella (1) Fomitopsis (5) Gloeophyllum (1) Gloeoporus (1) Gloeostereum (1) Grifola (2) Hirschioporus (3) Ischnoderma (1) Laetiporus (2) Laricifomes (1) Lenzites (1) Meripilus (1) Microporus (2) Oxyporus (1) Phaeolus (1) Piptoporus (1) Polyporus (10) Poria (1) Porodisculus (1) Pycnoporus (1) Rigidoporus (2) Trachyderma (1) Trametes (8) Trichaptum (1) Tyromyces (5)	70–90	70–100	Ohtsuka et al. 1973; Ito et al. 1976; Ohtsuka et al. 1977; Fujii et al. 1979; Liou and Lin 1979; Min et al. 1980; Nakajima et al. 1986; Kanayma et al. 1986; Mizuno et al. 1992; Gasiorowski et al. 1993; Cho et al. 1996;	
			Nanba 1998; Fullerton et al. 2000	
Schizophyllaceae – Schizophyllum(1)	70	-	Ohtsuka et al. 1973; Okamura et al. 1986	

Table 1 (continued)

Taxa (number of species studied)	A ctivity against:		Source	
	Sarcoma 180 solid cancer	Ehrlich solid cancer		
Gasteromycetideae				
Gasteromycetales				
Lycoperdaceae – Lycoperdon(2) Phallaceae – Dictyophora (1) Kobayasia (1) Boletales	-	-	Song et al. 1998 Miyazaki et al. 1975; Ukai et al. 1983; Hara et al. 1991; Ishiyama et al. 1996	
Boletaceae – Boletinus (1) Boletus (11) Filoboletus (1) Gyroporus (1) Leccinum(2) Phylloporus (1) Pulveroboletus(3) Suillus (5) Tylopilus (3)	70–100	90	Ohtsuka et al. 1973	
Xerocomus(3) Paxillaceae – Hygrophoropsis(1) Paxillus (3)	60-90	70–80	Ohtsuka et al. 1973	
Strobilomyceteceae –Boletellus (2) Porphyrellus (1) Strobilomyces(1)	60–80	60–70	Ohtsuka et al. 1973	
Gomphidiaceae –Gomphidius(1) Chroogomphus(1)	60–90	60–80	Ohtsuka et al. 1973	
Agaricomycetideae				
Agaricales				
Hygrophoraceae –Camarophyllus (2) Hygrocybe (14) Hygrophorus (21) Pleurotaceae –Pleurotus (4)	60–100	70–100 –	Ohtsuka et al. 1973 Yoshioka et al. 1972; Chung et al. 1982; Zhuang et al. 1994a; Song et al. 1998	
Tricholomataceae – Armillariella (3) Asterophora(1) Baeospora (1) Cantharellula (1) Catathelasma (2) Clitocybe (7) Collybia (6) Dictyopanus (1) Flammulina (1) Hohenbuehelia(1) Hypsizygus (1) Laccaria (6) Lampteromyces (1) Lepista (3) Leucopaxillus (1) Lyophyllum (8) Macrocystidia (2) Marasmiellus (2) Marasmius (6) Melanoleuca (2) Mycena (19) Omphalina (1) Oudemansiella (3) Panellus (1) Pleurocybella (1) Pseudohiatula (2) Resupinatus (1) Tricholoma (19 Tricholomopsis (4) Xeromphalina (3) Xerula (2)	60–100	60–100	Ohtsuka et al. 1973; Chung et al. 1982; Ikekawa et al. 1982; Kim et al. 1982; Ma et al. 1991; Ikekawa et al. 1992; Kiho et al. 1992a, b; Mizuno et al. 1994; Liu et al. 1996; Wang et al. 1996; Song et al. 1998; Ukawa et al. 2000	
Entolomataceae – Clitopilus (2) Entoloma (14) Rhodocybe (1) Rhodophyllus (6)	60-90	60-100	Ohtsuka et al. 1973	
Cortinariaceae – Cortinarius (25) Galerina (6) Gymnopilus(3) Hebeloma(3) Inocybe (19) Rozites (1)	60–100	60–100	Ohtsuka et al. 1973	
Bolbitiaceae – Agrocybe(7) Bolbitius (2) Conocybe(7)	60–90	70–90	Ohtsuka et al. 1973; Yoshida et al. 1996; Song et al. 1998	
Strophariaceae –Hypholoma(1) Kuehneromyces(1) Naematoloma(4) Pholiota (8) Psilocybe (3) Stropharia (2)	60–100	70–100	Ohtsuka et al. 1973; Chung et al. 1982; Song et al. 1998	
Crepidotaceae – Crepidotus (3) Tubaria (1)	60–100	90–100	Nakayoshi et al. 1968; Ohtsuka et al. 1973	
Amanitaceae – Amanita (21) Limacella (1)	60–100	60–90	Ohtsuka et al. 1973; Kiho et al. 1994; Yoshida et al. 1996	
Pluteaceae –Pluteus (5) Volvariella (4)	60–100	70–100	Ohtsuka et al. 1973; Chung et al. 1982; Misaki et al. 1986	
Agaricaceae – Agaricus (1) Cystoderma(2) Lepiota (15) Leucocoprinus(3) Macrolepiota (2) Melanophyllum(1) Phaeolepiota (1)	60–100	60–100	Ohtsuka et al. 1973; Mizuno 2002	
Coprinaceae – Coprinus (16) Panaeolus (1) Psathyrella (7) Pseudocoprinus(1) Russulales	60–100	60–100	Ohtsuka et al. 1973	
Russulaceae – Lactarius (18) Russula (23)	60-100	70-100	Ohtsuka et al. 1973	

Reshetnikov et al. 2001). Naturally collected or artificially growing fruit bodies, pure culture mycelia, and culture filtrate (culture broth) all contain biologically active polysaccharides.

Procedures for polysaccharide purification

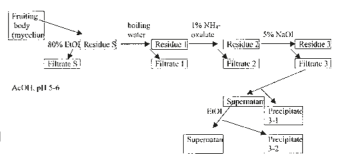
After two decades of intensive research on medicinal mushrooms, Mizuno and his co-workers in Japan developed reliable procedures for successful extraction, frac-Fig. 1 Fractional preparation of polysaccharides from mushrooms tionation and purification of polysaccharides from fruit- [adapted from Mizuno (1999a) with modification] ing bodies or culture mycelia. In general, this scheme involves elimination of low molecular weight substances from mushroom material using 80% ethanol, followed 2% ammonium oxalate (100°C, 6 h), and 5% sodium (1 3), (1 hydroxide (80°C, 6 h) (Mizuno 1996, 1999a).

acetic acid, ion-exchange chromatography, gel filtration, 1995b). and affinity chromatography. Basically, ion-exchange chromatography through DEAE-cellulose columns sepa- D-glucose upon acid hydrolysis (Mizuno 1996, 1999a). sorbed fraction) and β-glucans (non-absorbed fraction) ture (Marchessault et al. 1977). Acidic glucuronoxylowith the help of gel filtration and affinity chromatogra- mannan isolated from the fruit body of remella fucifor-(after elution with 1 M NaCl) yields purified polysaccha- fold helical backbone conformation (Yui et al. 1995). rides (Mizuno 1999a).

saccharides from mushrooms are shown schematically in structures have been described. These polysaccharides Fig. 1. It should be noted that the particular fractionation have linear or branched molecules in a backbone comprocedure scheme depends in each case on the polysac-posed of α - or β -linked glucose units, and they contain charide composition of the extracted material.

Structural composition of antitumor polysaccharides nent or in different combinations. in mushrooms

Polysaccharides belong to a structurally diverse class of sified as galactans, fucans, xylans, and mannans by the macromolecules, polymers of monosaccharide residues individual sugar components in the backbone. Heterojoined to each other by glycosidic linkages. It is note-glycan side chains contain arabinose, mannose, fucose, worthy that, in comparison with other biopolymers such galactose, xylose, glucuronic acid, and glucose as a main as proteins and nucleic acids, polysaccharides offer the component or in different combinations. highest capacity for carrying biological information because they have the greatest potential for structural vari-polysaccharides of different chemical structure from ability. The nucleotides in nucleic acids and the amino higher Basidiomycetes mushrooms has been investigatacids in proteins can interconnect in only one way ed; the main types are presented in Table 2. whereas the monosaccharide units in polysaccharides of branched or linear structures (Sharon and Lis 1993). can be seen in an analysis of polysaccharides of fruit bod-This enormous potential variability in polysaccharide ies of Pleurotus pulmonarius (= P. sajor-caju): 16 polyregulatory mechanisms of various cell-cell interactions different levels of antitumor activity (Zhuang et al. 1993, in higher organisms.



Mushroom polysaccharides are present mostly as gluby three successive extractions with water (100°C, 3 h), cans with different types of glycosidic linkages, such as 6)- β -glucans and (1 3)- α -glucans, but some are true heteroglycans. The others mostly bind to The first extraction results in water-soluble polysac- protein residues as PSP complexes (PSPC; Gorin and charides, the other two in water-insoluble polysaccha-Barreto-Berger 1983). The main source of antitumor rides. Polysaccharides extracted are further purified us- polysaccharides appears to be fungal cell walls that coning a combination of techniques, such as ethanol concen-sist of polysaccharides. However, chitin and chitosan tration, fractional precipitation, acidic precipitation with (fungal chitin) have no antitumor activity (Mizuno et al.

β-D-glucan is a polysaccharide yielding exclusively rates neutral polysaccharides from acidic ones. Neutral As for structure of schizophyllan tertiary conformation, polysaccharides are then separated intα-glucans (adactive β-D-glucan has a triple-strand right-winding strucphy. The same procedure with acidic polysaccharides mis was also demonstrated as having a left-handed, three-

Besides the well-known antitumorβ-(1 3)-glucans, General schemes for fractional preparations of poly- a wide range of biologically active glucans with other side chains that are attached in different ways. Heteroglucan side chains contain glucuronic acid, xylose, galactose, mannose, arabinose, or ribose as a main compo-

> Glycans, in general, are polysaccharides containing units other than glucose in their backbone. They are clas-

> A wide range of antitumor or immunostimulating

The number of antitumor active fractions in the fruit can interconnect at several points to form a wide variety bodies of mushrooms is remarkably high. One example structure gives the necessary flexibility to the precise saccharide fractions from 21 extractions demonstrated Table 3).

Table 2 Chemical structure of antitumor and immunostimulating polysaccharides of higher Basidiomycetes

t- Polysacchari	de	Species	R eferences	
Glucans				
α -(1 3)-glucan Linear α -(1 3)-glucan		Armillariella tabescens Amanita muscaria Agrocybe aegerita	Kiho et al. 1992a Kiho et al. 1994 Yoshida et al. 1996	
α-(1 4)-; β- α-(1 6)-; α- β-(1 6)-glu	-(1 4)- glucan	Agaricus blazei Agaricus blazei Lyophyllum decastes Armillariella tabescens	Fujimiya et al. 1998b Mizuno et al. 1990a Ukawa et al. 2000 Kiho et al. 1992a	
β-(1 6)-; β-	(1 3)-glucan	Agaricus blazei Grifola frondosa	Mizuno et al. 1990a	
β-(1 6)-; α- β-(1 3)-glu Mannoxylog Galactoxylog Xyloglucan	curonoglucan lucan	Agaricus blazei Ganoderma lucidum Grifola frondosa Hericium erinaceus Grifola frondosa Polyporus confluens	Nanba et al. 1987 Mizuno et al. 1990a Saito et al. 1989 Mizuno et al. 1986 Mizuno 1999b Mizuno et al. 1986 Mizuno et al. 1992	
Xylogalactog Mannogalac		Pleurotus pulmonarius Inonotus obliquus Pleurotus pulmonarius Pleurotus cornucopiae Ganoderma lucidum Agaricus blazei	Zhuang et al. 1993 Mizuno et al. 1999a Gutiérrez et al. 1996 Kim et al. 1994 Cho et al. 1999	
Galactomannoglucan		Flammulina velutipes Hohenbuehelia serotina Leucopaxillus giganteus	Ikekawa et al. 1982 Mizuno et al. 1994 Mizuno et al. 1995a	
Arabinogluc Riboglucan	an	Ganoderma tsugae Agaricus blazei	Zhang et al. 1994b Cho et al. 1999	
Glycans				
Arabinogalactar Glucogalactar α-(1 6)-ma Fucomannog Mannogalac Xylan Glucoxylan Mannogluco α-(1 3)-ma Glucomanna Glucomanna β-(1 2)-; β-	an n nnofucogalactan galactan tan tofucan oxylan nnan	Pleurotus citrinopileatus Ganoderma tsugae Sarcodon aspratus Fomitella fraxinea Dictyophora indusiata Pleurotus pulmonarius Grifola frondosa Hericium erinaceus Pleurotus pulmonarius Pleurotus pulmonarius Hericium erinaceus Dictyophora indusiata Agaricus blazei Agaricus blazei	Zhang et al. 1994a Wang et al. 1993 Mizuno 2000 Cho et al. 1998 Hara et al. 1991 Zhuang et al. 1993 Zhuang et al. 1994a Mizuno 1999b Mizuno 1999b Zhuang et al. 1993 Mizuno 1999b Ukai et al. 1983 Hikichi et al. 1999 Tsuchida et al. 2001 Mizuno et al. 1999b	
Galactogluco	omannan	Lentinus edodes	Fujii et al. 1979	

can with Man:Gal:Xyl:Glc in the polysaccharide at a mogalactan consisting of Xyl:Man:Gal (9:35:56 molar ratio). rides are FII-1 protein-containing xylan; FIII-1a proteincontaining glucoxylan consisting of Glc:Xyl (40:44 molar ratio), and FIII-2a protein-containing xyloglucan consisting of Xyl:Glc (36:62 molar ratio).

Correlation of structure and antitumor activities of mushroom polysaccharides

their chemical composition and configuration, as well as such as heteroß-glucans (Mizuno et al. 1995b), heterotheir physical properties. Antitumor activity is exhibited glycan (Gao et al. 1996b), β-glucan-protein (Kawagishi

The most antitumor-active water-soluble fractions by a wide range of glycans extending from homopolyfrom P. pulmonarius are Fi_o-a protein-containing xyloglu- mers to highly complex heteropolymers (Ooi and Liu 1999). Differences in activity can be correlated with lar ratio 2:12:42:42, and FA-2 protein-containing manno- solubility in water, size of the molecules, branching rate and form. Although it is difficult to correlate the struc-The most antitumor-active water-insoluble polysaccha- ture and antitumor activity of complex polysaccharides, some relationships can be inferred.

It is obvious that structural features such a \S -(1 3) linkages in the main chain of the glucan and additional β -(1 6) branch points are needed for antitumor action. β-glucans containing mainly (1 6) linkages have less activity. High molecular weight glucans appear to be more effective than those of low molecular weight (Mizuno 1996, 1999a, b). However, obvious variations in antitumor polysaccharides have also been noted. Antitu-Polysaccharides with antitumor action differ greatly in mor polysaccharides may have other chemical structures,

Table 3 Structure and antitumor activity of Pleurotus pulmonarius fruit bodies polysaccharides against Sarcoma 180 in mice (after Zhuang et al. 1993). FI-FA Water-soluble, FII-FIII water-insoluble polysaccharides

Polysaccharid MW ×	MW $\times 10^3$		Total sugar (%)	Component sugar (molar %)				Tumor inhibition
		(%)		Glc	Xyl	Man	Gal	ratio at 3 weeks (%)
FI _o -a	278	24.1	75.6	43.7	42.3	1.9	11.8	84.8
Fl _o -a-α	420	23.5	69.5	24.1	72.5	2.7	0.7	53.1
Fl _o -a-β	68	26.3	67.0	53.5	27.2	1.6	17.7	49.8
Fl _o -b-α	10	42.1	52.6	56.0	40.7	3.3	_	59.4
Fl _o -b-β	24	6.9	84.6	_	16.2	_	83.8	31.7
FA-1	11	27.5	67.7	71.6	5.5	_	22.9	48.7
FA-2	115	16.2	76.1	_	9.4	34.6	56.0	74.6
FA-3	10	75.3	22.5	50.0	14.9	13.1	22.0	34.5
FII-1	19	20.5	62.2	5.2	91.2	_	3.6	90.8
FII-2	17	44.1	50.5	9.2	86.2	_	4.6	8.0
FII-3	13	49.0	50.1	2.9	80.5	_	16.6	8.4
FIII-1a	87	70.5	15.4	39.8	43.7	7.8	8.7	76.9
FIII-1b	24	96.8	3.0	_	97.9	_	2.1	51.6
FIII-2	627	2.8	69.6	33.9	40.3	_	1.9	84.5
FIII-2a	700	2.5	68.8	62.2	35.5	_	2.3	100.0
FIII-2b	190	4.5	74.8	30.9	69.1	_	_	84.6

et al. 1990), α -manno β -glucan (Mizuno et al. 1995b), ous influence on the activity of the heteroglycans (Gao α-glucan-protein (Mizuno et al. 1995b) and heteroglycan- et al. 1996b). protein complexes (Zhuang et al. 1993; Mizuno et al.

A triple-helical tertiary conformation of medicinal Activation of mushroom polysaccharides mushroom β -(1 3)-glucans is known to be important by chemical modification for their immune-stimulating activity. When lentinan was denatured with dimethyl sulfoxide, urea, or sodium Different approaches to improving antitumor activity of et al. 1988). The same results, which confirm the corre- polysaccharides have been developed foGanoderma lation between antitumor activity and triple helix struc-lucidum, Grifola frondosa and Leucopaxillus giganteus ture, were obtained upon investigation of schizophyllan (= Tricholoma gigantea). These schemes include two (Yanaki et al. 1983, 1986).

many of these activities, such as macrophage nitrogen 1996, 1999a; Mizuno et al. 1996). Five polyaldehydes are independent of this conformation, e.g., synthesis of ously obtained from G. frondosa liquid culture myceli-2000), thus indicating that the α -(1 3)-mannan back- were first oxidised to polyaldehydes by 0.1 M NalO₄ in structure of the molecule.

that are strongly dependent on high molecular weight; ure (Zhuang et al. 1994b). Chemical activation of mushmushrooms, are not strongly dependent on molecular99% HCOOH solution; the reaction solution is then preacidic hydrolysate fractions of T. fuciformis fruit bodies is lyophilized after dialysis, while the other part is disof from 53 to 1 kDa that induce human monocytes to pro-alcohol precipitation (Zhuang et al. 1994b). Four fordue to the common structure of the (1 3)-mannan four polysaccharide fractions obtained from front front front four polysaccharide fractions obtained from front fr

hydroxide, tertiary structure was lost while primary mushroom polysaccharides by chemical modification structure was not affected, but tumor inhibition proper-have been described in the literature. The most successties were lowered with progressive denaturation (Maedaful schemes for chemical improvement of mushroom main procedures: modification of mushroom polysaccha-Mushroom β-(1 3)-glucans exhibit a variety of bio- rides by Smith degradation (oxydo-reducto-hydrolysis) logical and immuno-pharmacological activities, and and activation by the method of formolysis (Mizuno oxide synthesis, and limulus factor G activation, are de- and ten polyalcohols were prepared by the Smith degrapendent on the triple-helix conformation, while others dation method from five polysaccharide fractions previinterferony and colony stimulating factor (Yadomae um. For this reason, original polysaccharide solutions bone structure is of more importance than the tertiarydarkness, then converted into polyalcohols by reduction of NaBH 4 in alkaline medium adjusted to pH 8 with 2 M

Unlike β -(1 3)-glucans with medicinal properties NaOH, and hydrolysed by 1 M H₂SO₄ at room temperaranging from 500 to 2,000 kDa (Mizuno 1996), α-(1 3)- room polysaccharides by the method of formolysis inglucuronoxylomannans, which are characteristic of Jelly volves degradation of polysaccharides by formic acid in weight. Thus, Gao and co-workers (1996a) reported that cipitated with 99% EtOH, and one-half of the precipitate contain glucuronoxylomannans with molecular weights solved in hot water and additional fractions obtained by duce interleukin-6 as efficiently as non-hydrolyzed het- mylated polysaccharides and four formolysis products eropolysaccharide. This indicates that the activity may be of polysaccharides were prepared by this method from backbone; differences in molecular weight had no obvi-liquid culture mycelium. Although two of the original

polysaccharides had no activity, their polyaldehyde poly- Schizophyllum communéltoh et al. 1990; Hirata et al. ol, formylated, and formolysis derivatives showed signif- 1994). It was suggested that the sulfur content in schizprepared from a polysaccharide with low antitumor ac- man immunodeficiency virus (HIV) than the molecular charide (Zhuang et al. 1994b). As all original polysac- 1990; Hobbs 1995). The medicinal tests indicate that sulcharide fractions showing elevated activity levels by fated schizophyllan with a sulfur content of 5% can be chemical modification were β-glucan or xyloglucan, it useful as an anti-HIV agent for treatment of HIV-infectwas suggested that the sugar chain was changed or elimi-ed hemophiliacs (Hirata et al. 1994; Hobbs 1995). nated upon treatment, resulting in improved solubility It is important to realize that chemical modification is and activity (Mizuno 1999a).

used to transformβ-glucans into a water-soluble form. clinical qualities, most importantly water solubility and For example, whole fruit bodies of Pleurotus ostreatus the ability to permeate stomach walls after oral ingesor their stipes homogenate were treated with 0.15 M so-tion. dium hydroxide solution at 95°C for 2 h. The residue collected was washed with water until neutral, then suspended in 0.06% sodium chlorite solution, adjusted to Testing antitumor and immunomodulating activity pH 4.5 with acetic acid, and stirred for 6 h at 50°C. of mushroom polysaccharides The polysaccharide obtained wasβ-(1 3)-linked glucan, with every fourth glucopyranosyl residue substi- Initial data on the antitumor activity of mushroom extuted at 0-6 with singlep-glucopyranosyl groups. The tracts was circumstantial and in no way solid and reliheterogeneous etherification of the particulate glucanable. However, many indirect data that were properly medium gave the sodium salt of the water-soluble eficial effects of mushrooms on human health. A good O-(carboxymethyl) glucan derivative (Kuniak et al. example is an epidemiological study in Nagano Prefec-1993; Karácsonyi and Kuniak 1994). Carboxymethylated ture, Japan, where activity was monitored for several demodulatory effects, especially increased phagocytic rate of farmers whose main occupation was producing activity (Paulik et al. 1996).

linear α-(1 3)-glucan obtained from fruiting bodies of general population in the Prefecture (Ikekawa 1995, Amanita muscaria and Agrocybe aegeritahad little or no 2001). Another similar observation in Brazil brought antitumor effect, while their carboxymethylated products about extensive studies – and popularity – oAgaricus showed potent antitumor activity (Kiho et al. 1994; blazei (see below). Yoshida et al. 1996).

with amylase, cellulase, and protease (Kosuna 1998).

tained after enzymatic reduction of side chains and pro-mushroom, Hypsizygus marmoreus (Ikekawa 2001). ry and anticancer properties (Ghoneum et al. 1995; H. marmoreus. All mice were i.p. injected with a strong cal resection (Kidd 2000).

rally occur in higher Basidiomycetes mushrooms. Chem- 2001). ically sulfated schizophyllans with different sulfur con-

icant activity. Polyaldehyde, and polyol-polysaccharides ophyllan is more important in inhibiting growth of hutivity showed activity higher than the original polysac- weight or the nature of the sugar component (Itoh et al.

necessary in many cases to improve not only the antitu-Carboxymethylation is the other chemical method mor activity of mushroom polysaccharides, but also their

with monochloroacetic acid (GH₃ClO₂) in alkaline collected and processed gave good evidence for the benglucan from P. ostreatus (pleuran) exhibited immuno- cades. Researchers demonstrated that the cancer death Flammulina velutipes (a well known medicinal mush-In a similar manner, a water-insoluble, alkali-soluble room in Japan) was remarkably lower than that of the

I would like to emphasize the principal points of anti-Chemical modification of branched mushroom poly- tumor and immunomodulating effects of mushroom polysaccharides resulting in side-chain reduction can be de-saccharides. Most important among them are: (1) prevenveloped not only by Smith degradation but also by enzy-tion of oncogenesis by oral consumption of mushrooms matic reactions. A novel linear polysaccharide compris- or their preparations; (2) direct antitumor activity against 4)-bonded α-D-glucose units of a molecular various allogeneic and syngeneic tumors; (3) immunopoweight of 500-10,000 kDa was developed after succes- tentiation activity against tumors in combination with sive enzymatic treatments of submerged culture brothchemotherapy; (4) preventive effect on tumor metastasis.

A good example of preventive effect is given by Japa-Linear low molecular weight α -(1 4)-glucans ob- nese research on their popular edible and medicinal tein component (active hexose correlated compounds -Control mice were bred on an ordinary diet and treated AHCC) were demonstrated as having immunomodulato- mice with a diet containing 5% dried fruit body of Matsushita et al. 1998). In 1992, a trial was done in carcinogen, methyl-cholanthrene, and carcinogenesis of Japan to evaluate the preventive effect of AHCC against the mice was investigated. At the end of the 76-week obrecurrence of hepatocellular carcinoma following surgi- servation, 21 of the 36 control mice developed tumors, but only 3 of 36 mice in the treated group had tumors. Sulfated homo- and heteropolysaccharides possessingThe authors concluded that the mechanism of cancerantiviral activity are widespread in algae, especially in inhibitory and cancer-preventing activities of edible sea algae (Schaeffer and Krylov 2000), but do not natu- mushrooms was due to immunopotentiation (Ikekawa

It is well known from clinical practice that mushroom tent were obtained fromβ-(1 3)-glucan produced by polysaccharides work best in conjunction with other forms of 'tough' chemotherapy and surgery, which is, nan became a widely used medicine and dietary suppleunfortunately, very invasive and has a lot of negative ment in Japan, other Far East countries, and later in the side effects. Lentinan has been studied best in this re- United States and Europe. spect, both in animal models and in human clinical practice. In one study, 275 patients with advanced or recur- mune-enhancing activity and a broad antineoplastic rent gastric cancer were given one of two kinds of che-scope. It has been shown to prolong the survival time of motherapy (mitomycin C with 5-fluorouracil or tegafur) radiated mice, stimulate phagocytotic activity of macroeither alone or with lentinan injections. The best results phages, and improve the functions of the reticuloendowere obtained when lentinan was administered prior tothelial system (Zhu 1987). With regard to its antitumor chemotherapy and in patients with a primary lesion who properties, it acts directly on tumor cells, as well as indihad undergone no previous chemotherapy. The resultsrectly in the host to boost cellular immunity (Hobbs were evaluated on the basis of prolongation of life, re- 1995; Stamets 2000). It has shown antitumor activity in gression of tumors or lesions, and the improvement of animals with adenosarcoma, fibrosarcoma, mastocytoimmune responses (Hamuro and Chihara 1985; Hobbs ma, plasmacytoma, melanoma, sarcoma, carcinoma, and 1995; Wasser and Weis 1997a).

cancer therapy. A preventive effect of mushroom ex- jection of PSK at one tumor site has been shown to intracts on cancer metastasis has been studied by manyhibit tumor growth at other sites, thus helping to prevent groups, especially at the National Cancer Center Re- metastasis. PSK has been used both orally and intravesearch Institute of Japan. In a successful series of experi- nously in clinical medicine. It has been shown to be ments, Lewis lung carcinoma was s.c. transplanted into effective against many types of cancer (Hobbs 1995; the foot pads of mice and EA6 or EA6-PII (polysaccharides from Flammulina velutipe) were p.o. administered administered alone. for a period of 10 days. The life span of the group treated with EA6-PII was significantly increased (Ikekawa 2001). Further study was carried out using Meth-A fibro- 180, as well as against the solid form only of sarcoma 37, sarcoma: 7 days after the tumor was s.c. transplanted in- Erlich sarcoma, Yoshida sarcoma and Lewis lung carcito the abdomen of female BALB/c mice, the solid tumor noma (Hobbs 1995). Schizophyllan has also increased of each mouse was surgically dissected out, and 7 days cellular immunity by restoring suppressed killer-cell after the surgery, a second challenge with the same tu-activity to normal levels in mice with tumors (Borchers mor, Meth-A fibrosarcoma was made s.c. into the other et al. 1999). Best results against radiation damage were tumor growth was observed. The results indicated that or at the same time as radiation, and schizophyllan repre-treatment with EA6 slightly inhibited growth of the stored mitosis of bone marrow cells previously suprechallenged tumor, but post-treatment was remarkablypressed by anticancer drugs (Zhu 1987). Human clinical 10 mg/kg (lkekawa 2001).

rooms have undergone a lot of testing in animal model ex-vical carcinoma (Hobbs 1995). periments and in clinics. After isolation of lentinan from Lentinus edodesby Chihara in 1969, most of the experimental antitumor testing was performed with this polysac-Mechanisms of antitumor and immunomodulating charide. Its 'father', Chihara himself, was one of the first action by mushroom polysaccharides researchers to report the antitumor properties of lentinan.

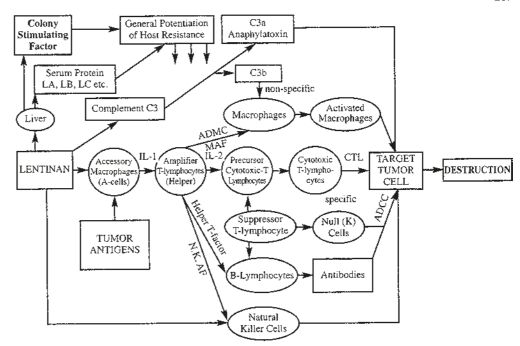
Originally, its effect was tested by using Sarcoma 180 Mushroom polysaccharides exert their antitumor action transplanted in CD-1/ICD mice (Chihara et al. 1969, mostly via activation of the immune response of the host 1970). Later, lentinan showed prominent antitumor ac-organism. These substances are regarded as biological tivity not only against allogenic tumors, but also against response modifiers (BRMs; Wasser and Weis 1999). This various synergic and autochtonous tumors (Hamuro andbasically means that: (1) they cause no harm and place Chihara 1985). Injections of lentinan into mice produced no additional stress on the body; (2) they help the body either an 80% reduction in tumor size or complete regres- to adapt to various environmental and biological stresssion in most of the animals tested (Chihara 1981). A numery es; and (3) they exert a nonspecific action on the body, ber of clinical tests followed. Among the first of these was supporting some or all of the major systems, including a follow-up, randomized control study on Phase 3 patients nervous, hormonal, and immune systems, as well as regwith advanced and recurrent stomach cancers (Wasser andulatory functions (Brekhman 1980).

Weis 1999; Ikekawa 2001). Lentinan therapy showed very The immunomodulating action of mushroom polysacgood results in prolonging the life span of patients and hadcharides is especially valuable as a prophylactic, a mild no toxic side effects. Similar results were obtained in pa- and non-invasive form of treatment, and in the preventients with colorectal and breast cancers. Since then, lenti-tion of metastatic tumors, etc., as described above. Poly-

PSK (commercial name krestin) has remarkable immammary, colon, and lung cancer (Sugimachi et al. Metastasis is a very serious and important problem in 1997). An intriguing feature of this compound is that in-Stamets 2000), but seldom with satisfactory results if

The polysaccharide schizophyllan shows antitumor activity against both the solid and ascite forms of Sarcoma side of the abdomen of the mouse and the re-challenge found when schizophyllan was administered shortly after effective for tumor growth inhibition at a dose of studies proved the beneficial activity of treatment with schizophyllan for patients with recurrent and inoperable Specific preparations from particular medicinal mush- gastric cancer, stage 2 cervical cancer, and advanced cer-

Fig. 2 Possible pathways of lentinan action (after Chihara 1981)



saccharides from mushrooms do not attack cancer cells tion of cytokines in peritoneal macrophages. Lentinan directly, but produce their antitumor effects by activating also increases peritoneal macrophage cytotoxicity different immune responses in the host. This has been against metastatic tumors; it can activate the normal and verified in many experiments, such as the loss of the alternative pathways of the complement system and can antitumor effect of polysaccharides in neonatal thymec-split C3 into C3a and C3b, enhancing macrophage actitomized mice or after administration of anti-lymphocyte vation (Aoki 1984; Wasser and Weis 1997a; Hobbs serum (Ooi and Liu 1999). Such results suggest that the 2000).

antitumor action of polysaccharides requires an intact Lentinan's immune-activating ability may be linked T-cell component and that the activity is mediated with its modulation of hormonal factors, which are through a thymus-dependent immune mechanism. Also,known to play a role in tumor growth. Aoki (1984) the antitumor activity of lentinan and other polysaccha-showed that the antitumor activity of lentinan is strongly rides is inhibited by pretreatment with antimacrophagereduced by administration of thyroxin or hydrocortisone. agents (such as carrageenan). Thus, the various effects of Lentinan can also restore tumor-specific antigen-directed polysaccharides are thought to be due to potentiation oflelayed-type hypersensitivity reaction.

the response of precursor T cells and macrophages to Schizophyllan activates macrophages (in vitro and in cytokines produced by lymphocytes after specific recog- vivo), which results in augmentation of T-cell activities nition of tumor cells (Hamuro and Chihara 1985). In and increases sensitivity of cytotoxic LAK and NK cells addition, the induction of a marked increase in the to IL-2 (Mizuno 1996). Although structurally related to amounts of CSF, IL-1, and IL-3 by polysaccharides relentinan, schizophyllan does not directly activate T-cells sults in maturation, differentiation, and proliferation of (Hobbs 1995). Possible pathways of such actions for the immunocompetent cells for host defense mechanismsentinan have been summarized in Chihara (1981) and (Hamuro and Chihara 1985). Mushroom polysaccharides Hamuro and Chihara 1985), and reviewed by Wasser and are known to stimulate natural killer cells, T-cells, Weis (1999), and those for β-p-glucan BRMs (Mizuno B-cells, and macrophage-dependent immune system2002) are shown in Figs. 2 and 3.

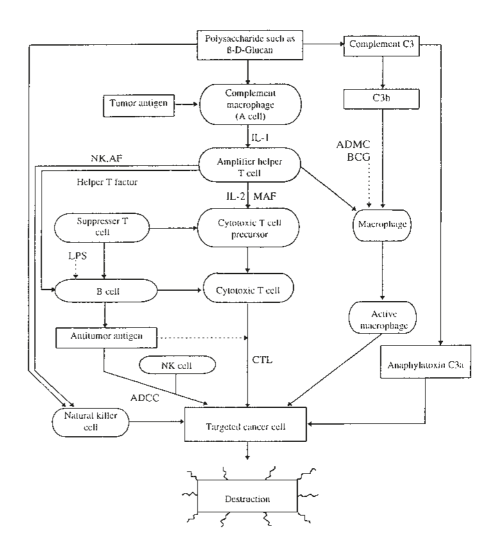
Lentinan is known to be able to restore the suppressed_activity of helper T-cells in the tumor-bearing host to Selected examples ofimportant medicinal their normal state, leading to complete restoration of hu-mushrooms with antitumor polysaccharides moral immune responses (Ooi and Liu 1999). The same in fruit bodies and cultured mycelium effect is true for PSK, while it has no substantial effect

on immune responses of the host under normal condi-One of the utmost important edible and medicinal biotions.

technological species, known as A. blazei was revaluated Infiltration of eosinophils, neutrophils, and granu- by our group. Analysis of data on cultivated mushroom

Infiltration of eosinophils, neutrophils, and granu- by our group. Analysis of data on cultivated mushroom locytes around target tissues is also accelerated by lenti- originating from Brazil, and study of type material of nan. It activates secretion of active oxygen and produc- A. blazei Murrill shows dramatic differences between

Fig. 3 Possible immune mechanism: β-D-glucan biological response modifier (BRM) (after Mizuno 2002)



1994; Ebina and Fujimiya 1998; Fujimiya et al. 1998a,

them. On the basis of existing differences the correct ma tumor-bearing mice (Kawagishi et al. 1989, 1990; name for widely cultivated mushroom was proposed as Mizuno et al. 1990b, 1998; Mizuno 2002; Itoh et al. new for science species Agaricus brasiliensis S. Wasser et al. A. blazei is the North American endemic not culti- 2000; Stamets 2000). Of the 17 polysaccharide fractions vated species known from only three localities – one in obtained from A. blazei fruit bodies (Mizuno et al. 1990a;

Florida and two in South Carolina (Wasser et al. 2002). Mizuno 2002), 7 were demonstrated to have antitumor Agaricus blazei mushroom (the Royal Sun Agaricus, activity. Analyses of physico-chemical properties of ABM, Himematsutake, Cogmelo de Dues) is one of the water-soluble polysaccharide fractions having high antimore newly discovered medicinal mushrooms. This deli-tumor activity showed that their main components were cious edible mushroom is native to a very small area of β -(1 6)-; β -(1 3)-glucan, acidic β -(1 6)-; α -(1 4)the mountains of Brazil, near the town of Sao Paulo. Epi-glucan, and acidic β -(1 6)-; α -(1 3)-glucan (Mizuno et demiologists studying the native population in this areaal. 1990a). A. blazei was the first mushroom described to found that they had a very low incidence of a variety of contain antitumor glucan with $\alpha\beta$ -(1 illnesses, including cancers as well as viral and bacteria- bone, unlike the well-knownβ-(1 3)-glucans. The antiinduced diseases, and a disproportionately high numbertumor proteoglucan HM3-G from A. blazei fruit bodies, enjoyed longevity. Eventually this was correlated with which mediated natural killer cell activation and apoptoconstant consumption of A. blazei mushroom in their nor-sis, has a molecular mass of 380 kDa and consists mal diet. During the 1980s and 1990s,A. blazei was dem- of more than 90% glucose, the main component being onstrated to be an immune system stimulant, promoting α -(1 4)-glucan with β -(1 6)-branching, at a ratio of the body's natural defense mechanisms to fight a variety approximately 4:1 (Fujimiya et al. 1998b). It is interestof infectious agents and conditions, including cancer. The ing to note that a low molecular weight fraction, LM-3, immunostimulating activity and antitumor action of with an average weight of 20 kDa, composed of A. blazei extracts were investigated in different laboratory α -(1 4)-glucan with β -(1 6)-branching, also demonmodels, including Sarcoma 180 and (Meth-A) fibrosarco- strated tumor-specific cytocidal and immunopotentiating 63.6:17.6:12.7, respectively; and AG-6 was composed of glucose and ribose at a molar ratio of 81.4:12.6.

A xyloglucan (Xyl:Glc, molar ratio =2:10) containing 9% protein obtained by fractionation and purification β -(1 of A. blazei extract showed significant activity against et al. 1986; Jong and Birmingham 1990; Wasser and Sarcoma 180 in mice (Mizuno 2002).

A. blazei are a source of antitumor polysaccharides. An Water-insoluble fractions include an acidic xyloglucan antitumor organic substance called ATOM was devel- with a Glc:Xyl molar ratio of 100:82 and 16.5% glucuoped from A. blazei (Iwade strain 101), which is a PSPC (Ito et al. 1997). Another PSPC, 0041, was obtained from submerged culture mycelium; the main components and three acidic glycoproteins with molecular masses of of this polysaccharide are glucose and mannose (Hikichi 20–100 kDa. The major component sugar is glucose, et al. 1999). A new antitumor polysaccharide active while fucose, xylose, mannose and galactose are minor against Sarcoma 180 was recently separated from liquid components (Mizuno et al. 1986). Thus, all polysacchacultured mycelium of A. blazei: β-(1 2)-; β-(1 3)-glucomannan (Tsuchida et al. 2001). This polysaccharide with different chain conformation, heteroglucans, or gluappears to be completely different from the antitumorcoproteins. polysaccharides from fruiting bodies of A. blazei (Mizuno et al. 1999b).

activity (Mizuno 2002).

structures. Polysaccharides from fruit bodies represented found (Zhuang et al. 1994a). Thus, polysaccharides from or heteroglucans; culture mycelia contained glucoman-xylans, or their complexes with protein, i.e., types of polyculture medium under submerged cultivation.

Ganoderma tsugaeis the other medicinal mushroom both the fruit body and mycelia. Seven glycans with nutrient medium used for cultivation. Thus, Ohno and costrong antitumor activities were obtained from 14 water-workers (1985, 1986) concluded that the antitumor glucan soluble and 15 water-insoluble fractions extracted from grifolan extracted from cultured mycelium of f. frondosa G. tsugae fruit bodies (Wang et al. 1993). Water-soluble is a β -(1 3)-, β -(1 6)-glucan, the same as in the fruit ated with mannose and fucose, and water-insoluble frac-was growing in liquid medium in stationary culture or tions represented protein-containing-(1 3)-glucans with shaking. The mycelium obtained was additionally with different protein content.

from G. tsugae mycelium and examined for antitumor β -(1 6)-glucans were obtained both by extraction of my-The three active polysaccharides obtained were: a gly-cipitation of buffer supernatant (Adachi et al. 2002).

effects (Fujimiya et al. 1999), whereas pure glucan ob- can-protein complex containing 9.3% protein, with a tained from antitumorβ-(1 6)-glucan-protein complex, heteropolysaccharide composed of mannose and xylose; isolated from water-insoluble residue of A. blazei fruiting a glucan-protein complex containing 25.8% protein; and bodies, did not exhibit strong activity (Kawagishi et al. a glycan-protein with glucose as the main component, 1990). Three immunostimulating heteroglucans (AG-2, and associated with arabinose, mannose, xylose, and -3, and -6) were extracted with 0.9% sodium chloride galactose. Comparison of active water-soluble polysacand hot water from fruiting bodies of A. blazei from charides obtained from fruit body and mycelium showed among the six polysaccharides obtained (Cho et al. 1999). that those from the fruiting body were glucogalactan-AG-2 and AG-3 were composed of glucose, galactose protein complexes, but those of the mycelium were and mannose in the molar ratios 74.0:15.3:10.7 and homoglucan-protein complexes or a heteroglycan composed of mannose and xylose.

Grifola frondosa is one of the most popular medicinal mushrooms. Fruit bodies of this mushroom contain 3)-; β -(1 6)-glucan, acidic β -D-glucan (Mizuno Weis 1999), and β -(1 6)-; β -(1 3)-glucan (Nanba et Not only fruit bodies but also cultured mycelia of al. 1987) in the water-soluble polysaccharide fraction. ronic acid; an acidic heteroglycan containing 3.8% protein, component sugars Glc:Xyl:Man:Fuc (100:58:34:14); rides detected inG. frondosa fruit bodies are β-glucans

In contrast to fruit body polysaccharide composition, no β-glucan has been detected among antitumor active A liquid medium filtrate separated from mycelium fractions obtained from culture mycelium (grown on after submerged cultivation of A. blazei contained man- Whatman filter paper soaked with liquid nutrient medium) nan-protein complex (AB-FP) with a molecular weight that was collected before initiation of fruit bodies (Mizuno of 10⁵–10⁷ Da and a small amount of glucose, galactose, and Zhuang 1995). In the water-soluble fractions, a fuand ribose. The yield of AB-FP was 575 mg/1 liquid cogalactomannan-protein complex, a glucogalactomanmedium filtrate, and it possesses significant antitumor nan, a mannogalactofucan, and a galactoglucomannofucan-protein complex were found. In water-insoluble frac-Thus, antitumor polysaccharides investigated in tions, a mannofucoglucoxylan, a mannoglucofucoxylan-A. blazei fruit body, culture mycelia, or produced extra- protein complex, a mannofucoglucoxylan-protein comcellularly in a culture medium have different chemical plex, and a glucomannofucoxylan-protein complex were glucans with different types of glucose unit connections G. frondosa are heteromannans, heterofucans, and heteronans, and mannan-protein complex was produced in asaccharide that were not found in fruit bodies of this mushroom.

It must be stated that the polysaccharide structure in in which polysaccharides have been well investigated in cultured mycelia may depend on the composition of the fractions were protein-containing glucogalactans associ-body of the mushroom. In this experiment, a pure culture cultivated for 3 days in a buffer composed of glucose Sixteen water-soluble polysaccharides were extracted (5%) and citric acid, pH 4.5. Antitumor active β -(1 3)-, effects on Sarcoma 180 in mice (Zhang et al. 1994b). celium grown on a nutrient medium and by alcohol pre-

Table 4 Number of polysaccharide fractions obtained from different Basidiomycetes

Species	Fruit body	Culture mycelium	R eferences
Agaricus blazei	17		Mizuno et al. 1990a
Hericium erinaceus	15		Mizuno 1999b
Grifola frondosa	29	28	Mizuno et al. 1986; Cun et al. 1994; Zhuang et al. 1994a
Hohenbuehelia serotina	20		Ma et al. 1991
Pleurotus pulmonarius	21		Zhuang et al. 1993
Pleurotus citrinopileatus	21		Zhang et al. 1994a
Leucopaxillus giganteus	24		Mizuno et al. 1995a
Lyophyllum decastes	11		Ukawa et al. 2000
Inonotus obliquus	21	8	Mizuno et al. 1999a
Ganoderma tsugae	29	16 ^a	Wang et al. 1993; Zhang et al. 1994b

^a Number of fractions of water-soluble polysaccharides only

The number of polysaccharides extracted from fruit- Table 5 Yield of polysaccharide fractions from sclerotia and culing body or cultured mycelium of the same species is ture mycelium of Inonotus obliquus(after Mizuno et al. 1999b) strongly dependent on the method of fractionation use but, in general, the total amount of polysaccharides in fruiting bodies is higher (Table 4).

The number of fractions indicated in Table 4 includes, in some cases, not only finally purified polysaccharides but also some intermediate fractions that were tested fo antitumor activity.

The proportion of biologically active polysaccharide fractions in fruit body and culture mycelium is very high. Thus, 20 of 29 polysaccharide fractions obtained from G. frondosa fruit body exhibited different levels of antitumor activity (Mizuno et al. 1986), and 24 of 28 polysaccharide fractions obtained from culture mycelium of this mushroom showed antitumor activity (Zhuang et al. 1994a).

the fruit body is higher, in general, than that obtained lecular weight, degree of branching, and higher (tertiary) water-soluble and water-insoluble polysaccharides ob- β -(1 3) linkages in the main chain of the glucan and tained from I. obliquus sclerotium is 2–3 times higher additional β -(1 6) branch points are needed for antituthan that extracted from cultured mycelium (Table 5).

Conclusions

ing thoroughly studied; even the inventory of known strongly dependent on molecular weight. species is incomplete, comprising maybe only 10% of the true number of species existing (Hawksworth 2001; activity of mushroom polysaccharides by chemical modi-Kirk et al. 2001). The number of mushrooms with fication, which is also necessary to improve their clinical Nevertheless, the species studied so far represent a vastwalls after oral ingestion. Two main procedures for Hetero- and Homobasidiomycetes, the overwhelming developed for Ganoderma lucidum Grifola frondosa and majority have been demonstrated to possess pharmaco-Leucopaxillus giganteus (= Tricholoma gigantea). Carture mycelia, or culture broth (Reshetnikov et al. 2001). forms β-glucans into a water-soluble form.

Mushroom polysaccharides are of different chemical

Water-soluble g/kg dry weig	polysaccharide ht	s, Water-inso g/kg dry v	Water-insoluble polysaccharides g/kg dry weight		
Sclerotium					
FIS-I	164.5	FII	2.64		
rFIS-II	12.0	FIII-1	42.48		
21		FIII-2	87.84		
Mycelium					
FI	53.9	FII	43.15		
		FIII-1	4.6		
		FIII-2	21.1		
-					

(Mizuno 1999a, 2000). The antitumor polysaccharides The total number of polysaccharides extracted from from various mushrooms are characterized by their mofrom culture mycelium. For example, the total of both structure. It is evident that such structural features as mor action. The β -glucans containing mainly (1 6) linkages have less activity. High molecular weight glucans appear to be more effective than those of low molecular weight (Mizuno 1996, 1999a, b). Unlike 3)-glucans, α-(1 3)-glucuronoxylomannans, β -(1 Higher Basidiomycetes mushrooms are still far from be- which are characteristic of jelly mushrooms, are not

Different approaches exist to improve the antitumor known pharmacological qualities is much lower still. qualities, water solubility and ability to permeate stomach source of anticancer and immunostimulating polysaccha-chemical improvement are: modification of mushroom rides. Many, if not all, Basidiomycetes mushrooms contain biologically active polysaccharides. Of the 651 spehydrolysis) and activation by the method of formolysis. cies and 7 infraspecific taxa from 182 genera of higher The most successful schemes for such methods have been logically active polysaccharides in their fruit bodies, cul- boxymethylation is another chemical method that trans-

A large body of experimental and clinical evidence composition, mainly belonging to the group of glucans demonstrates the beneficial results of mushroom polysaccharides for the following purposes: (1) prevention of Chihara G, Maeda Y, Hamuro J, Sasaki T, Fumiko F (1969) oncogenesis by oral consumption of mushrooms or their preparations; (2) direct antitumor activity against various Chihara G, Hamuro J, Maeda YY, Arai Y, Fukuoka F (1970) Fracallogeneic and syngeneic tumors; (3) immunopotentiation activity against tumors in conjunction with chemotherapy; (4) preventive effects on tumor metastasis. Most of the clinical evidence comes from the commercial polysaccharides lentinan, PSK (krestin), and schizophyllan, but there are also impressive new data for polysaccharides from Phellinus linteus, Flammulina velutipes Hypsizygus marmoreus A. blazei and others.

The biochemical mechanisms that mediate the biolog-Cho SM, Park JS, Kim KP, Cha DY, Kim HM, Yoo ID (1999)

I activity of polysaccharides are still not clearly un
Chemical features and purification of immunostimulating ical activity of polysaccharides are still not clearly understood. Polysaccharides from mushrooms do not attack cancer cells directly, but produce their antitumor effects

Korean J Mycol 27:170–174

Chung KS, Choi EC, Kim BK, Kim YS, Park YK (1982) The constituents and culture of Korean Basidiomycetes: anti-The antitumor action of polysaccharides requires an intact T-cell component; their activity is mediated through a thymus-dependent immune mechanism (Borchers et al. Cun Z, Mizuno T, Ito H, Shimura K, Sumiya T, Kawade M (1994) 1999). Mushroom polysaccharides are known to stimulate natural killer cells, T-cells, B-cells, and macrophagedependent immune system responses. The immunomod Ebina T, Fujimiya Y (1998) Antitumor effect of a peptide-glucan ulating action of mushroom polysaccharides is especially valuable as a means of prophylaxis, a mild and non-invasive form of treatment, prevention of metastatic tumors, and as a co-treatment with chemotherapy.

A wide range of biologically active polysaccharides is found among higher Basidiomycetes mushrooms, and their practical application is dependent not only on their unique properties but also on biotechnological availabili- Fujimiya Y, Suzuki Y, Oshiman KI, Kobori H, Moriguchi K, ty. Isolation and purification of polysaccharides from mushroom material is relatively simple and straightforward, and can be carried out with minimal effort (Mizuno 1996, 1999a). Mycelia formed by growing pure cultures in submerged conditions are of constantFujimiya Y, Suzuki Y, Katakura R, Ebina T (1999) Tumor-specific composition, and submerged culture is the best technique for obtaining consistent and safe mushroom products (Wasser et al. 2000; Reshetnikov et al. 2001).

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