

Wood Science and Technology

Green biorefinery of larch wood biomass with obtaining of bioactive compounds, functional polymers and nanoporous materials --Manuscript Draft--

Manuscript Number:	
Full Title:	Green biorefinery of larch wood biomass with obtaining of bioactive compounds, functional polymers and nanoporous materials
Article Type:	Original Article
Corresponding Author:	Boris Nikolaevich Kuznetsov, D.Sc. ICCT SB RAS Krasnoyarsk, RUSSIAN FEDERATION
Corresponding Author Secondary Information:	
Corresponding Author's Institution:	ICCT SB RAS
Corresponding Author's Secondary Institution:	
First Author:	Boris Nikolaevich Kuznetsov, D.Sc.
First Author Secondary Information:	
Order of Authors:	Boris Nikolaevich Kuznetsov, D.Sc. Irina G. Sudakova, Ph.D Natalya V. Garyntseva, Ph.D Vladimir A. Levdansky, D.Sc Natalya M. Ivanchenko, Ph.D Andrey V. Pestunov Laurent Djakovitch, D.Sc Catherine Pinel, D. Sc
Order of Authors Secondary Information:	
Author Comments:	
Suggested Reviewers:	Dmitry Murzin, D Sc. Abo Akademi University, Turku, Finland dmurzin@abo.fi Expert in the area of biomass conversial Paresh Dhepe, D Sc. National Chemical Laboratory, Pune, India pl.dhepe@ncl.res.in Expert in lignin valorization Oxana Taran, D Sc. Boreskov Institute of Catalysis, Novosibirsk, Russia oxanap@catalysis.ru Expert in area of biomass catalytic conversions Zinaida Pai, D Sc. Boreskov Institute of Catalysis, Novosibirsk, Russia zpai@catalysis.ru Expert in the area of catalytic valorization of biomass

[Click here to view linked References](#)

Green biorefinery of larch wood biomass with obtaining of bioactive compounds, functional polymers and nanoporous materials

B.N. Kuznetsov^{1,2}, I.G. Sudakova¹, N.V. Garyntseva¹, V.A. Levdansky¹, N.M. Ivanchenko¹, A.V. Pestunov¹,
L Djakovitch³, C. Pinel³

¹Institute of Chemistry and Chemical Technology SB RAS, FRC KSC SB RAS

Krasnoyarsk, 660036, Akademgorodok, 50-24, Russia

*e-mail: bnk@icct.ru, Fax/Tel +7(391)249-48-94

²Siberian Federal University, Krasnoyarsk, 660041, Svobodny prospect, 79, Russia

³IRCELYON, 2 avenue Albert Einstein, F-69626 Villeurbanne Cedex, Lyon, France

Abstract

The first green biorefinery of larch wood based on new methods of biomass fractionation on dihydroquercitin (DHQ), arabinogalactan (AG), microcrystalline cellulose (MCC) and soluble lignin (SL) is reported. The new green method of one-step isolation of DHQ and AG from larch wood by ethanol-water solution is described. The first results of kinetic studies and optimization of the process of extracted larch wood peroxide fractionation on MCC and SL in acetic acid – water medium in the presence of green TiO₂ catalyst are presented. The products obtained from larch wood were characterized by FTIR, SEM, AFM, XRD and chemical methods. The scheme of larch wood biorefinery is suggested which integrates the described processes of biomass fractionation on DHQ, AG, MCC, SL and the yearly developed methods of polysaccharides processing to bioactive copper and amino acid – containing sulfated polymers along with the methods of nanoporous materials synthesis from lignin. All these methods use such non-toxic and less-toxic reagents, as water, ethanol, hydrogen peroxide, acetic acid, sulfamic acid, urea.

Introduction

Annual growth of plant biomass far exceeds the requirements of mankind for fuel and chemical products (Naik et al. 2010; Davis et al. 2014). Actually, from the components of plant biomass it is possible to obtain the entire range of products of modern petrochemical synthesis, as well as unique natural compounds, in particular physiologically active substances. Lignocellulosic biomass mainly consists of cellulose, hemicelluloses and lignin (Sjostrom 1993; Rowell 2012). These plant polymers are structured in plant cells in a complex way and the lignocellulosic biomass is quite resistant to chemical reagents and enzymes (Pandey 2015; Furkan et al. 2015). Therefore, in industrial pulp and hydrolysis processes the chemically aggressive and environmentally hazardous reagents, elevated temperatures and pressures are used (Sixta 2006).

Traditional catalysts for biomass transformation processes are corrosive and environmentally hazardous mineral acids. The problems of their use are associated with the occurrence of side reactions and the difficulties in separation of target products and catalyst from the reaction medium.

Current research trends in the development of new efficient processes for the conversion of lignocellulosic biomass to valuable chemicals, functional polymers and bioethanol are associated with the use of solid catalysts and "green" reagents (Fang et al. 2016; Serrano-Ruiz and Dumesic 2011; Bessone et al. 2014). To increase the biomass reactivity and the area of its contact with solid catalyst, as well as to reduce the diffusion limitations in heterogeneous catalytic processes of biomass conversion, the various activation methods (mechanical, mechanochemical, chemical, etc.) are used (Ramos 2003; Zeng et al. 2009). The principle possibility of application of solid acid and bifunctional catalysts for hydrolysis of cellulose in aqueous medium was shown (Hu et al. 2015;

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

Glozzi et al. 2014; Ishida et al. 2014; Liu et al. 2015). In recent year, studies are actively conducted on development the integrated technologies of biomass complex processing (Cherubini et al. 2009; Ed de Long et al. 2013; Zhang et al. 2016; Zhang et al. 2015).

Promising integrated processes for biomass conversion include, as a key step, its fractionation on cellulose, sugars and lignin, from which the other various chemicals are obtained (Dagle et al. 2016; Daijang and Chen 2014; Xu et al. 2016, Zhang et al. 2015).

The authors suggested to use one-stage catalytic processes of wood fractionation in the medium "hydrogen peroxide-acetic acid-water" (Kuznetsov et al. 2008; Kuznetsov et al. 2009; Kuznetsov et al. 2015) and in supercritical ethanol (Baryshnikov et al. 2014; Sharypov et al. 2015).

Since the wood species differ in their chemical composition, and therefore in the composition of the products of their processing, an important task is to develop the optimal green methods for transformation the biomass of the most common tree species to the valuable chemicals, biofuels and functional polymers.

Larch is the predominant coniferous species in the forests of Siberia, Canada and North America. The high density of the wood and the large amount of extractives complicate the use of larch wood in the pulp and hydrolysis industries. The most effective way of the use of larch biomass is its complex chemical processing with obtaining valuable physiologically active compounds and new functional polymers (Horhammer et al. 2014).

Most of the works published in recent years are devoted to the extractive isolation of bioactive dihydroquercetin and arabinogalactan from larch wood (Ravber et al. 2015; Troshina and Roshchin 2014; Ma et al. 2012).

The task of the present study is to develop a new approach to the complex environmetally safe processing of larch wood, based on the integration in a single technological cycle the new "green" methods of biomass components conversion to a wide range of demanded products.

Materials and methods

Air-dry sawdust (fraction 2-5 mm) of larch wood (*Larix sibirica Ledeb*) harvested in the forest area near Krasnoyarsk city was used as initial raw material. Wood composition (wt%): cellulose 41.2, lignin 28.1, hemicellulose 26.4 (including arabinogalactan 19.9), dihydroquercetin 2.0, ash 1.0.

Isolation of dihydroquercetin and arabinogalactan

The flask volume of 2 liters, equipped with a reflux condenser was charged with 100 g larch wood, poured 1.5l of water or 5–25% aqueous solution of ethanol (special purity, Sigma Tech, Russia) and heated on a steam bath for 2 hours. Then solution was separated from the wood and concentrated under vacuum on a rotary evaporator to 60–80ml, while stirring it diluted with 300 ml of ethanol. The precipitated AG was separated by filtration, washed on a filter with 40-50 ml of ethanol and dried at room temperature. The mother solution in the amount of 400–430 ml remaining after separation of AG, was concentrated under vacuum until complete removal of the solvent. Then to the flask with a dry residue was added 350–400 ml of 96% ethanol, clarified with activated carbon and heated on a steam bath for 10 minutes and filtered through a filter alumina oxide. The resulting solution was concentrated under vacuum on a rotary evaporator to completely remove ethanol. DHQ was obtained in the form of loose powder of light yellow color.

Peroxide fractionation of larch wood

Catalytic delignification of wood sawdust by H₂O₂ was carried out as in (Kuznetsov et al. 2013) using 250 cm³ glass reactor equipped with mechanical stirrer, reflux condenser and thermometer. Wood sawdust (10 g) was

1 placed in the glass reactor. Then, a mixture of glacial acetic acid, hydrogen peroxide (all of special purity, Sigma
2 Tech, Russia), distilled water and commercial TiO₂ (analytical purity, DuPont, USA) was added. The reaction
3 mixture was vigorously stirred (700 rpm) at selected temperature (70–100 °C) during 1–4 h. Composition of the
4 reaction mixture was varied in the following range: hydrogen peroxide 3–7 wt%, acetic acid 15–40 wt%,
5 liquid/wood ratio (LWR) 10–15. Concentration of TiO₂ catalyst was at 1 wt%. TiO₂ has an average particle size of
6 about 10 microns, rutile phase composition and BET surface area of 3 m²/g. When the reaction was completed, the
7 solid product was separated under vacuum using Buchner funnel and drying at 105 °C until constant weight.
8

9 The residual lignin content in cellulosic product was used to evaluate the delignification activity of TiO₂
10 catalyst. The cellulosic product yield was estimated by gravimetric method and calculated as follows: $Y = (m/m_0) \times$
11 100, where Y – yield of cellulosic product, wt%; m – weight of abs. dry cellulosic product, g; m₀ – weight of abs.
12 dry wood, g.
13

14 Analytical methods

15 The content of cellulose, hemicelluloses and lignin in solid products was defined chemical methods,
16 generally accepted in wood chemistry (Sjööstroöm and Alern 1999).
17

18 X-ray diffraction analysis of cellulose products was carried out on PANalyticalX'Pert Pro diffractometer using
19 Cu-Ka source ($\lambda = 0,154$ nm) in the 2θ range 5–70 ° and scanning step width of 0.01 °/scan. The samples of cellulose
20 were analyzed by the powder method in cuvette with 25 mm diameter. Crystallinity index (CI) was calculated from the
21 ratio of the height between the intensity of the crystalline peak and total intensity after subtraction of the background
22 signal (Park et al. 2010):
23

$$24 \text{ CI} = (I_{002} - I_{AM}) / (I_{002}),$$

25 where I_{002} – is the height of the 002 peak; I_{AM} – is the height of the minimum between the 002 and the 101 peaks.
26

27 Infrared spectroscopy analysis (FTIR) was carried out in transmission mode. Samples of cellulose (4 mg for
28 each) were prepared in pellet with matrix KBr. The spectra were recorded with Bruker Tensor – 27 in the
29 range 4000–400 cm⁻¹ wavelength. Spectral data were processed by the program OPUS/YR (version 2.2).
30

31 The electron images were obtained with scanning electron microscope TM-1000 HITACHI (Japan) with
32 accelerating potential 15 kV. Samples were coated on carbon support.
33

34 AFM images were obtained with scanning probing multimode microscope Solver P47 (NT-MDT,
35 Moscow). Transparent films of samples were prepared by evaporation of their water solutions on flat support made
36 from glass. The rate of scanning was 1–2 Hz and the number of points on scanning area 256 × 256 or 512 × 512.
37

38 **Results and discussion**

39 Extraction of bioactive dihydroquercetin and arabinogalactan from larch wood

40 Bioactive dihydroquercetin (DHQ) and arabinogalactan (AG) are increasingly used in medicine,
41 pharmaceutical and food industries (Babkin et al. 1997; Goellner et al. 2011). AG is a promising precursor for
42 synthesis of new bioactive polymers (Medvedeva et al. 2003).
43

44 Larch wood contains to 2,5 % wt. of flavonoids and to 20 % wt. polysaccharide arabinogalactan. The
45 content of DHQ (3,3',4',5,7 – pentaoxyflavone) reaches 90 % from the sum of flavonoids. AG is usually obtained
46 by water extraction of larch wood (Medvedeva et al. 2003). The known methods of DHQ isolation are based on
47 larch wood extraction by organic solvents, mainly by ethylacetate and acetone (Babkin et al. 2000; Kislitsin and
48 Mal'chikov 2009; Gavrilov 2009). Organic solvents extract together with DHQ resinous substance which
49 contaminate the dihydroquercetin. In particular, the extraction of larch wood by 80-86% ethanol significantly
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

contaminated DHQ with resinous substances and complicated and lengthy purification was required to obtain the pure product (Gavrilov 2009).

We developed the new green method of one-step isolation of DHQ and AG by water-ethanol extraction on larch wood. This method provides the simultaneous production of DHQ and AG, which are not contaminated with resinous substances.

The effect of ethanol concentration in water solution on the yield of DHQ and AG from larch wood was studied. It was found that the maximum yield of DHQ (1.81% wt.) corresponds to the ethanol concentration of 15% and it does not increase with further rise of the ethanol content in the solution (Fig. 1a). However, the yield of AG decreases with the growth of ethanol concentration in water over 10% (Fig.1b).

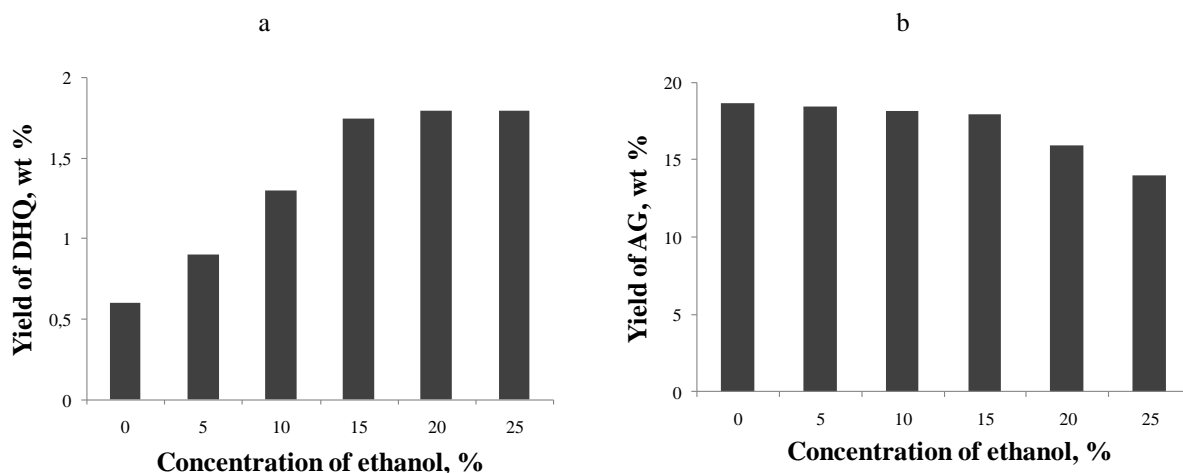


Fig.1 The yields of dihydroquercetin (a) and arabinogalactan (b) isolated from larch wood with ethanol-water solutions (extraction time 2 hours)

As follows from the obtained data, the optimum ethanol concentration which provides a high yield of DHQ (1.8 wt.%) and AG (18.0 wt%) corresponds to 15%.

It was established that a preliminary mechanical treatment of larch wood in a planetary mill for 2 min allows to shorten the time of the extraction of DHQ and AG from 2 to 0.5 hours without reducing their yields (Table.1).

Table 1 Influence of ethanol concentration on the yield and degree of extraction of DHQ and AG from larch wood, mechanically activated for 2 min*

Concentration of ethanol in water, %	Yield of DHQ from abs. dry wood, %	The degree of DHQ extraction, %	Yield of AG from abs. dry wood, %	The degree of AG extraction, %
0	0,52	26,0	18,7	94,0
5	0,83	41,5	18,5	93,0
10	1,3	65,0	18,3	92,0
15	1,81	90,5	18,0	90,4
20	1,78	89,0	16,9	84,9
25	1,82	91,0	15,1	76,0

*extraction time 0,5 h.

The structure of DHQ and AG isolated from larch wood were confirmed by FTIR and ¹³C NMR methods.

Peroxide catalytic fractionation of larch wood

1 The promising approaches for wood biorefinery include, as a key stage, woody biomass fractionation on
2 polysaccharides and lignin and their subsequent transformation to chemicals and liquid biofuels.

3 The known ways of lignocellulosic biomass fractionation can be divided into two groups. In one of them
4 the cellulose and hemicelluloses are removed and the lignin remains as a rest, another includes methods in which the
5 lignin is removed.
6

7 Reductive catalytic fractionation is the known way to separate lignocellulosic biomass into lignin-based
8 soluble mono-, di- and oligomers while retaining most of the carbohydrates in the pulp (Van den Bosch et al. 2015;
9 Schutyser et al. 2015; Anderson et al. 2016).
10

11 Catalytic oxidation of lignins by molecular oxygen in alkaline medium allows to separate lignocellulosic
12 biomass on aromatic aldehydes (mainly vanillin and syringaldehyde) and cellulose (Tarabanko et al. 2017).
13

14 Peroxide catalytic oxidation of hardwood and softwood in acetic acid – water medium enables to achieve
15 the effective separation of wood biomass components on microcrystalline cellulose and soluble products from lignin
16 and hemicelluloses (Kuznetsov et al 2008, 2009, 2015).
17

18 In earlier works, the authors proposed to carry out the catalytic fractionation of larch wood on cellulose and
19 soluble lignin in a "hydrogen peroxide-acetic acid-water" medium at temperatures of 120-130 ° C and increased
20 pressure in the presence of catalysts H₂SO₄, H₂MoO₂ and Fe₂(MoO₄)₃ (Kuznetsov et al. 2008; Kuznetsova et al.
21 2005; Kuznetsova et al 2005). In recent studies the authors demonstrated the possibility an effective catalytic
22 fractionation of aspen wood (Kuznetsov et al. 2013), birch wood (Garyntseva et al. 2015), fir wood (Kuznetsov et al.
23 2017) on microcrystalline cellulose and soluble lignin under mild conditions (temperature 100 ° C, atmospheric
24 pressure). This was achieved by reducing the contribution of diffusion limitations of the delignification process
25 when the hydromodule and the size of wood particles were decreased and the intensive mixing of the reaction
26 medium was used. The high catalytic activity in the peroxide delignification of birch wood and abies wood at mild
27 conditions demonstrated the suspended catalyst TiO₂ (Garyntseva et al. 2015; Kuznetsov et al. 2017).
28

29 In this study, the peroxide delignification process, catalyzed by TiO₂ was used for fractionation on MCC
30 and soluble products the larch wood from which the DHQ and AG were removed. The successful replacement the
31 dangerous H₂SO₄ catalyst and the expensive Mo catalyst on non-toxic and non-corrosive TiO₂ catalyst in peroxide
32 fractionation of extracted larch wood under mild conditions was demonstrated. The optimal process conditions
33 providing a high yield of cellulose with low content of residual lignin were determined.
34

35 To optimize the process of extracted larch wood peroxide fractionation in the presence of TiO₂ catalyst the
36 influence of temperature, concentrations of hydrogen peroxide and acetic acid, liquid/wood ratio, time on the
37 dynamics of lignin removal from wood was studied.
38

39 The increase of temperature from 70 °C to 100 °C, concentrations of H₂O₂ from 3 wt% to 6 wt%,
40 CH₃COOH from 15 wt% to 35 wt% and LWR from 5 to 15 reduces significantly the content of residual lignin and
41 of hemicelluloses in the cellulosic product (Table 2). But at the same time the yield of cellulosic product is
42 decreased. The oxidative destruction of lignin, hemicelluloses and the amorphous part of cellulose in wood is
43 accelerated when delignification temperature increased from 70 to 100 °C.
44

45 Delignification of larch wood at 70 °C during 4 h gives 60.2 wt% yield of cellulosic product containing
46 68.4 % of cellulose and 24.2 % of lignin. When raising the temperature to 100 °C, the yield of cellulosic product
47 decreased to 44.2 wt%. The maximum content of cellulose in the product (93.1 %) was detected after 4-hour
48 delignification of larch wood at 100 °C. Under such conditions, the residual lignin was actually absent in this sample
49 and hemicelluloses amount was 5.8 %.
50
51
52
53
54
55

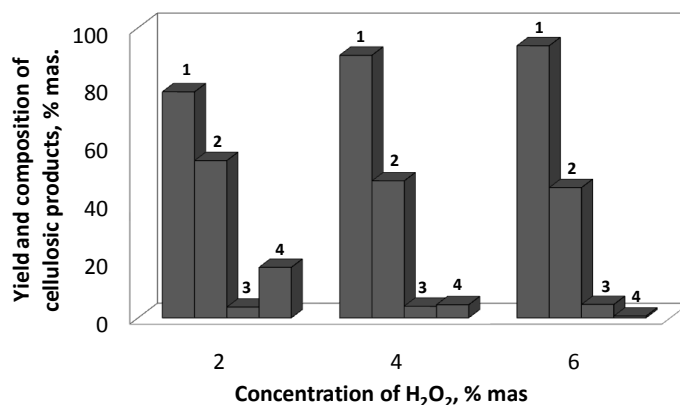
Table 2 Impact of delignification temperature on the yield and composition of cellulosic product obtained from extracted larch wood (CH₃COOH 25 wt%, H₂O₂ 5 wt%, LWR 15, TiO₂ 1 wt%)

Temperature, °C	Time, h	Yield of cellulosic product, wt%*	Composition of product, wt%**		
			Cellulose	Hemicellulose	Lignin
70	1	63,6	64,7	8,1	26,7
70	2	61,8	66,6	7,6	25,3
70	3	61,0	67,5	7,0	25
70	4	60,2	68,4	6,9	24,2
80	1	63,4	64,9	8,2	26,4
80	2	59,9	68,7	8,1	22,7
80	3	57,3	71,9	7,9	19,7
80	4	56,0	73,5	6,5	19,5
90	1	58,2	70,8	6,9	21,8
90	2	56,6	72,7	6,4	20,4
90	3	53,5	77,0	6,3	16,2
90	4	48,9	84,1	5,9	9,5
100	1	53,5	77,0	6,5	16
100	2	45,0	91,5	6,0	2
100	3	44,6	92,3	5,9	1,3
100	4	44,2	93,1	5,8	0,6

* on abs.dry wood, ** on abs.dry product

The influence of H₂O₂ concentration on cellulosic product yield and its composition was studied at 100 °C (Fig. 2). In the presence of 2.0 wt% of H₂O₂ the yield of cellulosic product is 77.9 wt%, but it has a rather high content of residual lignin (17.4 %). When the concentration of H₂O₂ increases, the content of residual lignin reduces, but the yield of cellulosic product is reduced simultaneously.

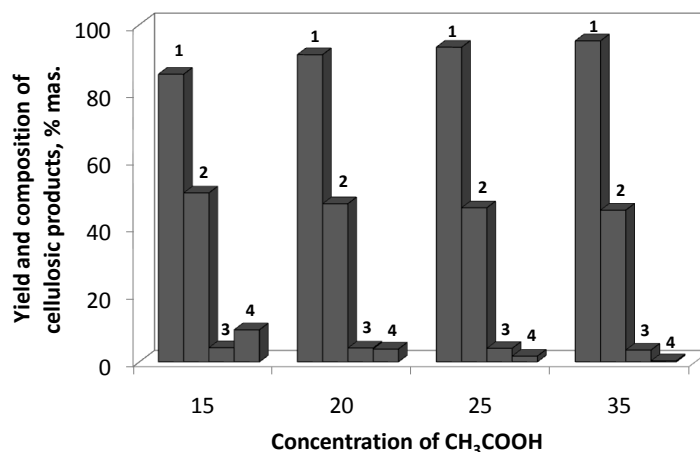
At the high H₂O₂ concentration, the oxidation of wood carbohydrates occurs along with the oxidation of lignin. According to the data obtained, the optimum concentration of H₂O₂, corresponding to the high yield of cellulosic product (near 45 wt% on a.d.w.) and low lignin content in the cellulosic product (0.6 – 4.5 wt% on a.d.p.) is 4–6 wt% (Fig. 2).



1 – cellulose, 2 – yield, 3 – hemicelluloses, 4 – lignin

Fig. 2 Impact of concentration of H₂O₂ on the yield and composition of cellulosic products obtained from extracted larch wood (CH₃COOH 30 wt%, 1 % TiO₂ catalyst, 100 °C, time 4 h)

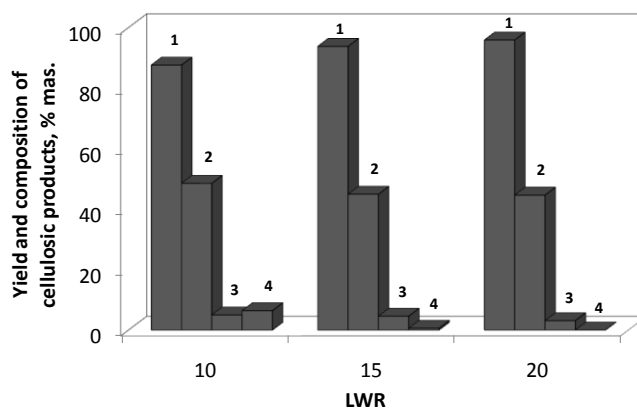
The increase of acetic acid concentration in reaction mixture decreases the yield of cellulosic product along with the reduction in residual lignin content (Fig. 3). While acetic acid concentration was increased from 15 to 35 wt%, the yield of cellulosic product decreased from 50.1 up to 45.0 %. Simultaneously, the content of cellulose in cellulosic product significantly increased from 85.3 up to 95.2 %. The concentration of acetic acid has also a strong influence on the residual lignin content. Its amount reached 9.5 % in the presence of 15 % wt of acetic acid and was lowered to 0.3 % when 35 % wt acetic acid was used. Experimental data presented in Fig. 3 show that the optimum concentration of acetic acid is nearly 25 wt% This concentration allows to reach the rather high yield of cellulosic product (45.7 wt%) with high amount of cellulose (93.3 %) and low content of residual lignin (1.7 wt%).



1 – cellulose, 2 – yield, 3 – hemicelluloses, 4 – lignin

Fig. 3 Impact of concentration of CH₃COOH on the yield and composition of cellulosic products obtained from extracted larch wood (H₂O₂ 6 wt%, 1 % TiO₂ catalyst, 100 °C, time 4 h)

The value of the liquid/wood ratio (LWR) also allows controlling both the yield and composition of cellulosic products (Fig. 4). At liquid/wood ratio of 15–20 they have low content of residual lignin. The reduction in LWR to 10 increases both the amount of cellulosic product (48.5 %) and residual lignin concentration (6.4 %). A possible reason is a hindered diffusion of lignin oxidation products from wood to solution at low LWR. As a result, the small molecules of lignin oxidative fragmentation are re-condensed to so-called pseudo lignin (Hu et al. 2012). According to obtained data an optimal LWR value for studied process of larch wood delignification is equal to 15.



1 – cellulose, 2 – yield, 3 – hemicelluloses, 4 – lignin

Fig. 4 Impact of LWR on the yield and composition of cellulosic products obtained from larch wood (H₂O₂ 6 wt%, CH₃COOH 30 wt%, 1 % TiO₂ catalyst, 100 °C, time 4 h)

The kinetic study of larch wood peroxide catalytic delignification in the temperature range 70–100 °C was accomplished. The variation of lignin concentration in the cellulosic product was used for calculating the rate constants of delignification process. It was found that the process of peroxide delignification of larch wood in the presence of TiO₂ catalyst is described by the first order equations (Fig.5).

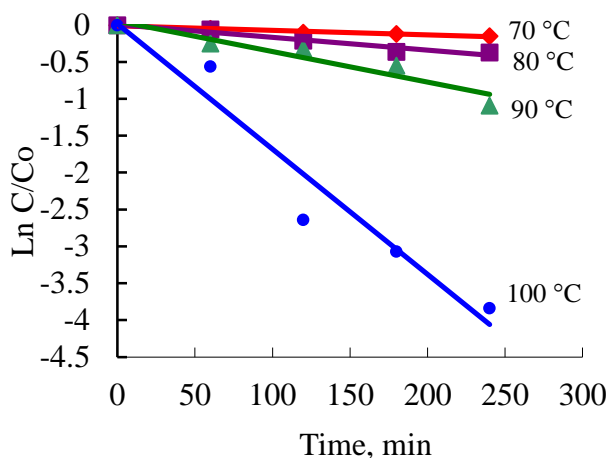


Fig.5 Dynamic of lignin isolation from extracted larch wood in the presence of 1 % TiO₂ catalyst (H₂O₂ 5 wt%, CH₃COOH 25 wt%, LWR 15)

The calculated rate constants of larch wood peroxide delignification are presented in Table 3.

Table 3 Rate constants of extracted larch wood peroxide delignification (H₂O₂ – 6 wt%, CH₃COOH – 30 wt%, catalyst TiO₂ – 1 wt%, LWR – 15)

Temperature, °C	$k \cdot 10^{-4}, \text{c}^{-1}$
70	0,10
80	0,28
90	0,68
100	2,84

The activation energy of larch wood peroxide delignification process was determined using temperature dependence of the rate constants in Arrhenius coordinates (Fig. 6). The rather high value of activation energy (88 kJ·mol⁻¹) points on the minor contribution of external diffusion limitations at the used conditions of larch wood peroxide delignification in the presence of TiO₂ catalyst.

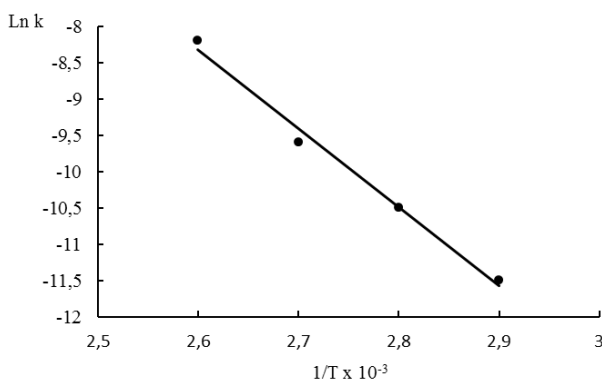


Fig. 6 Temperature dependence of rate constants of extracted larch-wood peroxide delignification (H₂O₂ 6 wt%, CH₃COOH 30 wt%, LWR 15, 1 wt% TiO₂)

The similar kinetic parameters of peroxide delignification of aspen wood, birch wood and abies wood (Kuznetsov et al. 2017) (the first order equations, the comparable rate constants and values of activation energy) point to the identical mechanism of delignification processes in the presence of H₂SO₄ and TiO₂ catalysts.

The numerical optimization of the process of larch wood peroxide delignification in the presence of TiO₂ catalyst was carried out with the use of Statgraphic application software, according to earlier described procedure (Sudakova et al. 2013). The main purpose of the analysis was to find conditions for the most complete removal of lignin from wood, while maintaining a sufficiently high yield of cellulosic product.

As independent parameters, the following factors have been selected: X₁ – the H₂O₂ concentration in reaction solution, wt%; X₂ – liquid to wood ratio. The other process parameters were fixed: temperature 100 °C, concentration of acetic acid 30wt %; TiO₂ 1 wt %, time 4 h.

The following output parameters for optimization were selected: Y₁ – the cellulosic product yield, wt%; Y₂ – the cellulose content in the product, wt%; Y₃ – lignin content in the product, wt%.

Optimization was performed with the use of generalized parameter of optimization (W_a) which was calculated using the following equation:

$$W_a = \frac{\sum_{j=1}^p \delta_j d_j}{\sum_{j=1}^p \delta_j}$$

where δ – the weight of the output parameter $0 \leq \delta_j \leq 1$; d_j – private utility function which was calculated using the following equation:

$$d_j = \frac{\varphi_0(x) - y_j^{(-)}}{y_j^{(+)} - y_j^{(-)}}$$

Where $\varphi_0(x)$ – the response of output parameter at the point X; $y_j^{(+)}$, $y_j^{(-)}$ – the best and worst values of the output parameters, within the studied region.

Table 4 presents the data on conditions and results of experiments for the calculation of generalized parameter of optimization. When choosing weights δ , it was assumed, that the residual lignin content is the most important parameter. For this reason this parameter got the weight equal 1. The weights of such parameters as yield of cellulosic product and cellulose content in the product were choosing equal 0.5.

Table 4 Matrix for planning of the experiments on larch wood delignification, the results of dispersion analysis, the utility functions and the generalized parameter of the optimization

Experimental number	X ₁	X ₂	Y ₁	Y ₂	Y ₃	d ₁	d ₂	d ₃	W _a
1	2	10	62,1	71,3	25,7	0,9526	0,0120	0,0280	0,2552
2	4	10	51,7	82,8	11,1	0,4053	0,4720	0,5560	0,4974
3	6	10	48,5	87,6	6,4	0,2368	0,6640	0,7440	0,5972
4	2	15	54,2	77,9	17,4	0,5368	0,2760	0,3040	0,3552
5	4	15	47,2	90,5	4,5	0,1684	0,7800	0,8200	0,7564
6	6	15	44,9	93,8	0,6	0,0474	0,9120	0,9760	0,7279
7	2	20	50,8	84,8	10,7	0,3578	0,5520	0,5720	0,5135
8	4	20	45,4	93,3	2,2	0,1556	0,8920	0,9120	0,6471
9	6	20	44,6	95,9	0	0,0316	0,9960	0,9989	0,7179
Y ⁽⁻⁾			44	71	25				
Y ⁽⁺⁾			63	96	0				
δ			0,5	0,5	1,0				

Table 5 shows the results of analysis on the influence of the main factors on the generalized parameter of optimization W_a.

Table 5 Analysis of variances for W_a.

Dispersion source	Sum of squares	df	F-Ratio	P-Value
A:X1	0,140791	1	26,87	0,0139
B:X2	0,0465873	1	8,89	0,0585
AA	0,0223943	1	4,27	0,1305
AB	0,00473344	1	0,90	0,4120
BB	0,011285	1	2,15	0,2385

Analysis of variances showed that the effect of both factors X_1 and X_2 on the generalized parameter of optimization is statistically significant (P-Value less than 0.05 and the confidence levels above 95%).

As a result of mathematical processing the following regression equation was obtained:

$$W_a = -1,1927 + 0,3398X_1 + 0,1215X_2 - 0,0264X_1^2 - 0,0034X_1X_2 - 0,0030X_2^2$$

Response surface of the generalized parameter of the optimization is presented on Fig. 7.

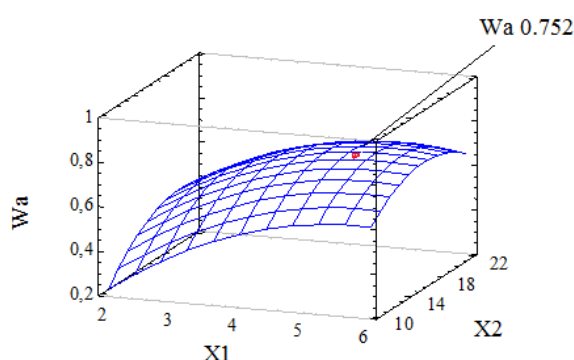


Fig. 7 Response surface of the generalized parameter (W_a) of the optimization of larch wood peroxide delignification with TiO_2 catalyst: X_1 – initial concentration of H_2O_2 , X_2 – liquid to wood ratio

It was found, that the generalized parameter of the optimization is set to 0.752. This corresponds to the following optimal parameters of larch wood delignification process: temperature 100 °C, concentrations of H_2O_2 6.0 % wt. and of CH_3COOH 30 % wt., LWR 15, duration 4 h.

Cellulosic product obtained by peroxide fractionation of extracted larch wood at optimal condition with an yield 44.2 wt% has the following composition (wt%): cellulose 93.1, hemicelluloses 5.8, lignin 0.6. According to FTIR, XRD and SEM data the structure of this cellulosic product corresponds to that of commercial microcrystalline cellulose (Fig. 8, 9, 10).

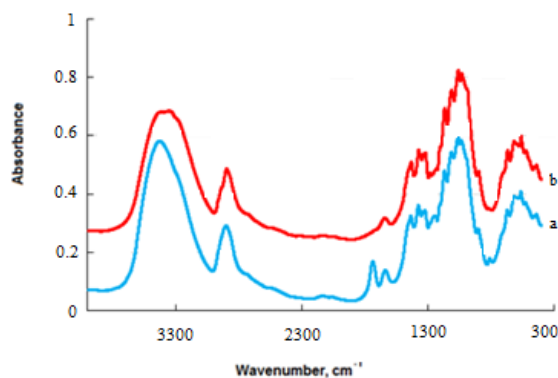


Fig. 8 FTIR spectra of cellulosic product from extracted larch wood (a) and of commercial microcrystalline cellulose Vivapur (b)

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65

In FTIR spectrum of cellulosic product from larch wood the adsorption band, which corresponds to either the acetyl or uronic ester groups of hemicelluloses (Xiang et al. 2016) appears in the region 1700–1740 cm^{-1} . This fact indicates the presence of residual hemicelluloses in the cellulosic product from extracted larch wood.

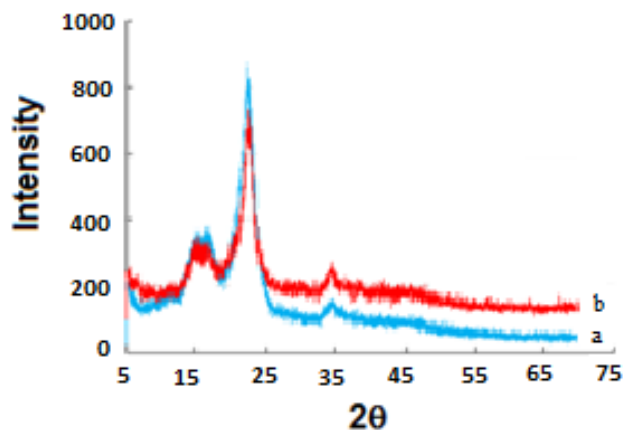


Fig. 9 Diffraction patterns of cellulosic product from extracted larch wood (a) and of commercial microcrystalline cellulose Vivapur (b)

Crystallinity index of cellulose from extracted larch wood, calculated as in paper (Park et al. 2010) is equal to 0.7.

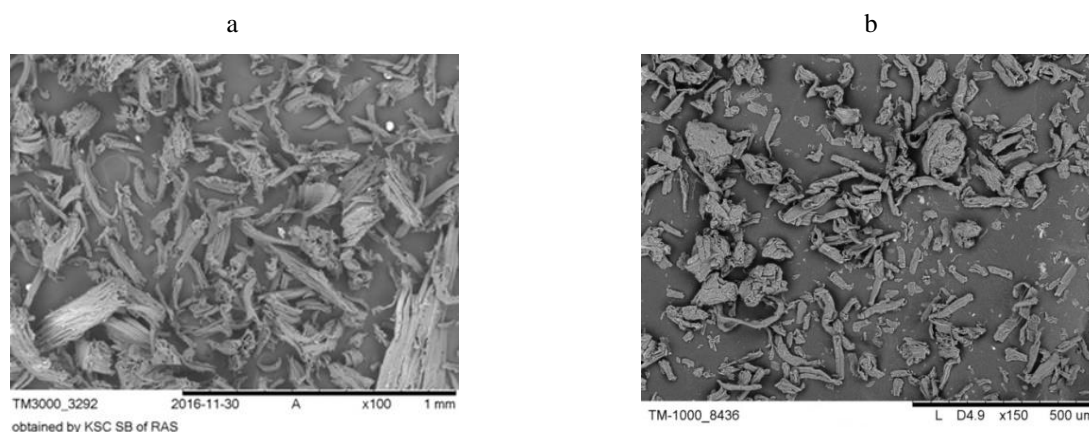


Fig 10 SEM images of cellulosic product from extracted larch wood (a) and of commercial MCC Vivapur 101 (b)

According to GC-MS data the soluble products of extracted larch wood peroxide fractionation contain only small amounts of aromatic compounds, indicating deep oxidation of phenylpropane units of lignin at studied process conditions. It is possible that the hydroxyl and peroxide radicals generated from H_2O_2 on the surface of TiO_2 catalyst can diffuse through the liquid reaction medium to the wood particles and to execute the oxidative destruction of lignin via homolytic mechanism (Kuznetsov et al. 2008).

Thus, the results of the study demonstrate the possibility of effective fractionation of extracted larch wood biomass into microcrystalline cellulose and soluble products from lignin under mild conditions (100°C, atmospheric pressure) in the medium “ H_2O_2 – CH_3COOH – H_2O – TiO_2 catalyst”.

Development of the scheme of green biorefinery of larch wood

In our previous papers (Kuznetsov et al. 2013; Kuznetsov et al. 2013) it was suggested to use as a key process of biorefinery of low-grade wood the peroxide catalytic fractionation of wood biomass on polysaccharides and lignin.

Based on this approach the scheme of green biorefinery of larch-wood biomass is proposed which integrates the optimized processes of biomass fractionation on DHQ, AG, MCC and soluble lignin. Polysaccharides and lignin are further processed with obtaining of bioactive sulfated polysaccharides and nanoporous materials (Fig.11).

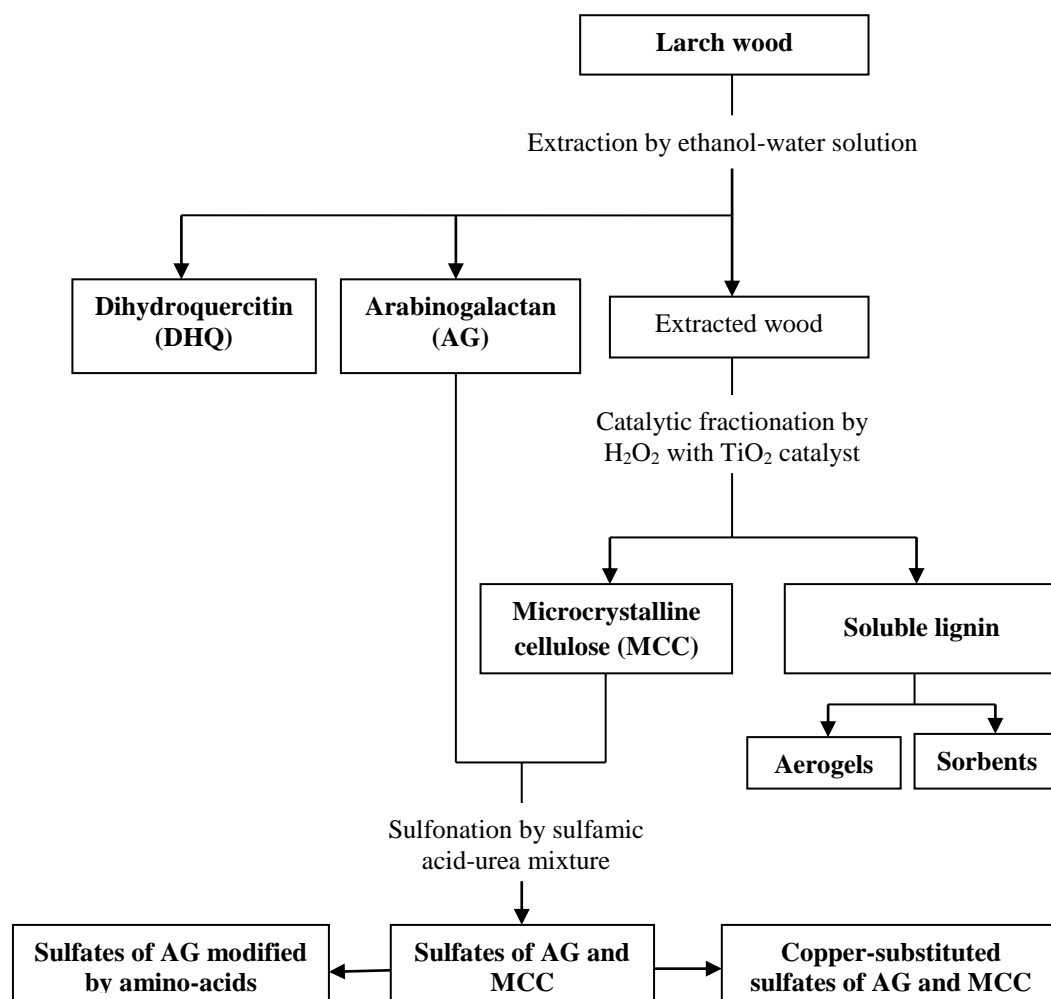
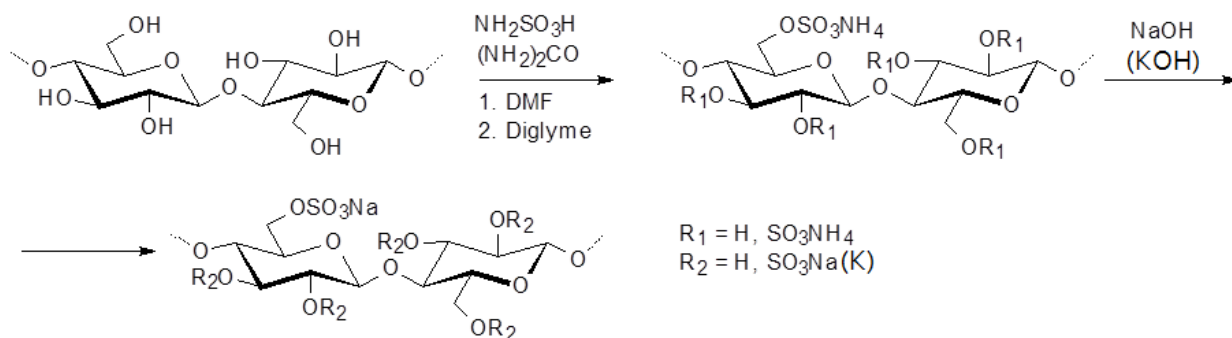


Fig. 11 Suggested scheme of green biorefinery of larch wood

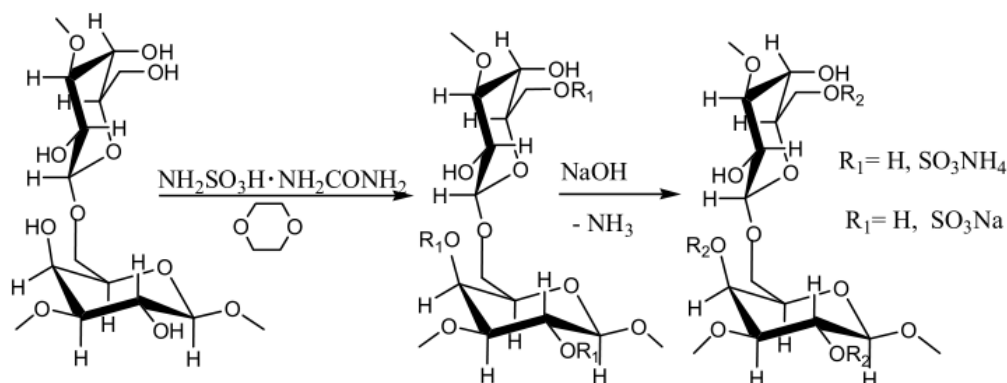
Described in this study new methods of DHQ and AG extraction from larch wood, catalytic peroxide fractionation the extracted wood on MCC and soluble lignin use such non-toxic agents, as water, ethanol, hydrogen peroxide, acetic acid.

For the synthesis of MCC and AG sulphates, we proposed yearly to use a low-toxic sulfamic acid and urea instead of dangerous and corrosive sulfating agents (Kuznetsov et al. 2015; Kuznetsov et al. 2017).

MCC sulfates are obtained as ammonium, sodium and potassium salts:



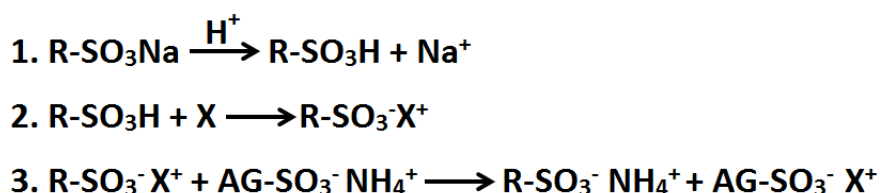
AG sulfates are obtained as ammonium and sodium salts (Kuznetsov et al. 2017):



Sulfated arabinogalactan can be used as water-soluble polymer for drug delivery.

On the basis of sulfated AG and MCC the new bioactive polymers, containing Cu^{2+} ions and amino-acids were synthesized by ion-exchange method.

The following scheme of obtaining the functional polymers of AG sulfates was used (Kuznetsov et al. 2017):



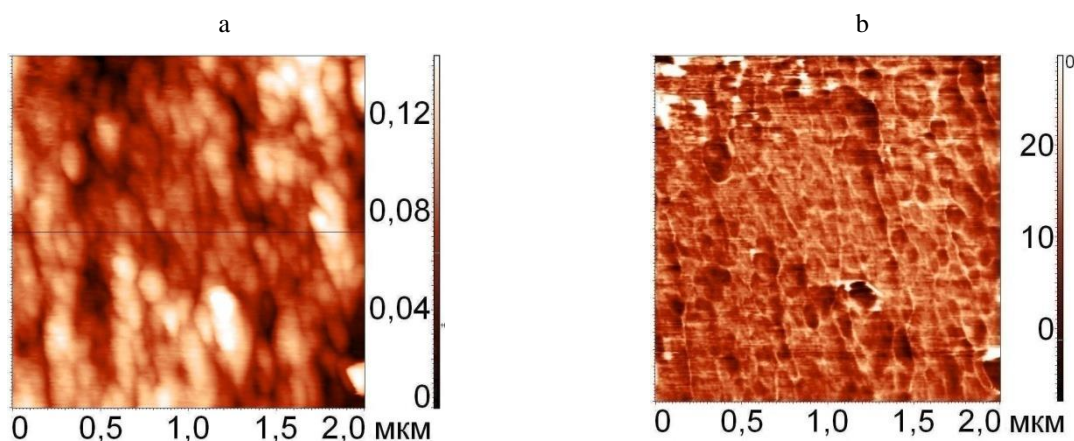
R – ion-exchange resin with $-\text{SO}_3\text{H}$ -group, X – Cu^{2+} , amino-acids: glycine, ornithine, histidine, arginine, AG – arabinogalactan.

Procedure of functional polymers synthesis included the following steps:

- Substitution of Na^+ on H^+ in the resin
- Sorption of Cu^{2+} ions or amino-acids on SO_3H -groups of resin.
- Ion exchange of NH_4^+ groups of sulfated AG on Cu^{2+} cations or on protonated amino-acid.

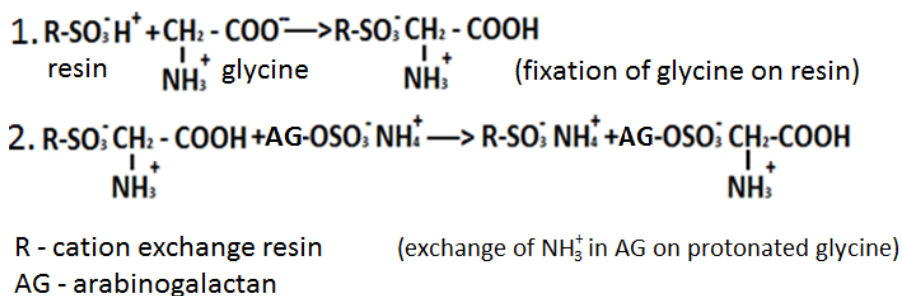
Copper-containing sulfates of AG are the water-soluble powders which form the transparent films with green color after water evaporation. Analysis of EPR spectra showed that these polymers contain the isolated and interacting Cu^{2+} ions (Kuznetsov et al. 2017). The concentration of isolated Cu^{2+} ions decreases with the growth of copper content in sulfated AG. It was proposed that the isolated Cu^{2+} ions are fixed on AG areas with low concentration of sulfate groups, but and they form salt-like structures on the AG areas with high concentration of sulfate groups.

1 Ion-exchange method was firstly used for synthesis of water-soluble copper-containing sulfates of
 2 microcrystalline cellulose (Cu-MCS). The copper content in the obtained Cu-MCS samples is 12.6-14.1%. The
 3 absence of sodium in the obtained polymer shows complete substitution of sodium cations on the copper cations in
 4 the sodium salt of MCC sulfate. The structure of the copper-containing sulfates of MCC was studied by XRD, FTIR,
 5 EPR and AFM methods. It was shown by XRD, that Cu-MCS and the sodium salt of MCC sulfate had amorphous
 6 structure in contrast to the samples of the original MCC, with a high degree of crystallinity. FTIR spectrum of
 7 copper-containing MCC sulfate was similar to spectrum of sodium salt of MCC sulfate. According to EPR data
 8 there is a formation of copper-containing salt system with a pseudo-crystalline structure in the samples of Cu-MCS.
 9 As shown by atomic-force microscopy, the surface of the Cu-MCS films consists of homogeneous crystallites which
 10 have a spherical or slightly elongated form with average size of about 70 nm (Fig. 12). Surface of the film is
 11 homogeneous and it contains no impurities.
 12
 13
 14



15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32 **Fig. 12** AFM images of film of copper-containing MCC sulfate with Cu content 14.1 wt% (a – relief, b – phase
 33 contrast)
 34
 35

36 The water-soluble derivatives of sulfated arabinogalactan with amino acids glycine, ornithine, histidine and
 37 arginine were prepared by ion exchange method (Vasilyeva et al. 2016; Vasilyeva et al. 2016). From the analysis of
 38 FTIR spectra it is concluded that the resulting compounds contain the negatively charged sulfated arabinogalactan as
 39 an anion and the protonated forms of amino-acids – as a cations. In particular, the following two-stage scheme was
 40 used for glycine fixation on sulfated AG (Vasilyeva et al. 2016):
 41
 42
 43
 44



45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65
 66
 67
 68
 69
 70
 71
 72
 73
 74
 75
 76
 77
 78
 79
 80
 81
 82
 83
 84
 85
 86
 87
 88
 89
 90
 91
 92
 93
 94
 95
 96
 97
 98
 99
 100
 101
 102
 103
 104
 105
 106
 107
 108
 109
 110
 111
 112
 113
 114
 115
 116
 117
 118
 119
 120
 121
 122
 123
 124
 125
 126
 127
 128
 129
 130
 131
 132
 133
 134
 135
 136
 137
 138
 139
 140
 141
 142
 143
 144
 145
 146
 147
 148
 149
 150
 151
 152
 153
 154
 155
 156
 157
 158
 159
 160
 161
 162
 163
 164
 165
 166
 167
 168
 169
 170
 171
 172
 173
 174
 175
 176
 177
 178
 179
 180
 181
 182
 183
 184
 185
 186
 187
 188
 189
 190
 191
 192
 193
 194
 195
 196
 197
 198
 199
 200
 201
 202
 203
 204
 205
 206
 207
 208
 209
 210
 211
 212
 213
 214
 215
 216
 217
 218
 219
 220
 221
 222
 223
 224
 225
 226
 227
 228
 229
 230
 231
 232
 233
 234
 235
 236
 237
 238
 239
 240
 241
 242
 243
 244
 245
 246
 247
 248
 249
 250
 251
 252
 253
 254
 255
 256
 257
 258
 259
 260
 261
 262
 263
 264
 265
 266
 267
 268
 269
 270
 271
 272
 273
 274
 275
 276
 277
 278
 279
 280
 281
 282
 283
 284
 285
 286
 287
 288
 289
 290
 291
 292
 293
 294
 295
 296
 297
 298
 299
 300
 301
 302
 303
 304
 305
 306
 307
 308
 309
 310
 311
 312
 313
 314
 315
 316
 317
 318
 319
 320
 321
 322
 323
 324
 325
 326
 327
 328
 329
 330
 331
 332
 333
 334
 335
 336
 337
 338
 339
 340
 341
 342
 343
 344
 345
 346
 347
 348
 349
 350
 351
 352
 353
 354
 355
 356
 357
 358
 359
 360
 361
 362
 363
 364
 365
 366
 367
 368
 369
 370
 371
 372
 373
 374
 375
 376
 377
 378
 379
 380
 381
 382
 383
 384
 385
 386
 387
 388
 389
 390
 391
 392
 393
 394
 395
 396
 397
 398
 399
 400
 401
 402
 403
 404
 405
 406
 407
 408
 409
 410
 411
 412
 413
 414
 415
 416
 417
 418
 419
 420
 421
 422
 423
 424
 425
 426
 427
 428
 429
 430
 431
 432
 433
 434
 435
 436
 437
 438
 439
 440
 441
 442
 443
 444
 445
 446
 447
 448
 449
 450
 451
 452
 453
 454
 455
 456
 457
 458
 459
 460
 461
 462
 463
 464
 465
 466
 467
 468
 469
 470
 471
 472
 473
 474
 475
 476
 477
 478
 479
 480
 481
 482
 483
 484
 485
 486
 487
 488
 489
 490
 491
 492
 493
 494
 495
 496
 497
 498
 499
 500
 501
 502
 503
 504
 505
 506
 507
 508
 509
 510
 511
 512
 513
 514
 515
 516
 517
 518
 519
 520
 521
 522
 523
 524
 525
 526
 527
 528
 529
 530
 531
 532
 533
 534
 535
 536
 537
 538
 539
 540
 541
 542
 543
 544
 545
 546
 547
 548
 549
 550
 551
 552
 553
 554
 555
 556
 557
 558
 559
 560
 561
 562
 563
 564
 565
 566
 567
 568
 569
 570
 571
 572
 573
 574
 575
 576
 577
 578
 579
 580
 581
 582
 583
 584
 585
 586
 587
 588
 589
 590
 591
 592
 593
 594
 595
 596
 597
 598
 599
 600
 601
 602
 603
 604
 605
 606
 607
 608
 609
 610
 611
 612
 613
 614
 615
 616
 617
 618
 619
 620
 621
 622
 623
 624
 625
 626
 627
 628
 629
 630
 631
 632
 633
 634
 635
 636
 637
 638
 639
 640
 641
 642
 643
 644
 645
 646
 647
 648
 649
 650
 651
 652
 653
 654
 655
 656
 657
 658
 659
 660
 661
 662
 663
 664
 665
 666
 667
 668
 669
 670
 671
 672
 673
 674
 675
 676
 677
 678
 679
 680
 681
 682
 683
 684
 685
 686
 687
 688
 689
 690
 691
 692
 693
 694
 695
 696
 697
 698
 699
 700
 701
 702
 703
 704
 705
 706
 707
 708
 709
 710
 711
 712
 713
 714
 715
 716
 717
 718
 719
 720
 721
 722
 723
 724
 725
 726
 727
 728
 729
 730
 731
 732
 733
 734
 735
 736
 737
 738
 739
 740
 741
 742
 743
 744
 745
 746
 747
 748
 749
 750
 751
 752
 753
 754
 755
 756
 757
 758
 759
 760
 761
 762
 763
 764
 765
 766
 767
 768
 769
 770
 771
 772
 773
 774
 775
 776
 777
 778
 779
 780
 781
 782
 783
 784
 785
 786
 787
 788
 789
 790
 791
 792
 793
 794
 795
 796
 797
 798
 799
 800
 801
 802
 803
 804
 805
 806
 807
 808
 809
 810
 811
 812
 813
 814
 815
 816
 817
 818
 819
 820
 821
 822
 823
 824
 825
 826
 827
 828
 829
 830
 831
 832
 833
 834
 835
 836
 837
 838
 839
 840
 841
 842
 843
 844
 845
 846
 847
 848
 849
 850
 851
 852
 853
 854
 855
 856
 857
 858
 859
 860
 861
 862
 863
 864
 865
 866
 867
 868
 869
 870
 871
 872
 873
 874
 875
 876
 877
 878
 879
 880
 881
 882
 883
 884
 885
 886
 887
 888
 889
 890
 891
 892
 893
 894
 895
 896
 897
 898
 899
 900
 901
 902
 903
 904
 905
 906
 907
 908
 909
 910
 911
 912
 913
 914
 915
 916
 917
 918
 919
 920
 921
 922
 923
 924
 925
 926
 927
 928
 929
 930
 931
 932
 933
 934
 935
 936
 937
 938
 939
 940
 941
 942
 943
 944
 945
 946
 947
 948
 949
 950
 951
 952
 953
 954
 955
 956
 957
 958
 959
 960
 961
 962
 963
 964
 965
 966
 967
 968
 969
 970
 971
 972
 973
 974
 975
 976
 977
 978
 979
 980
 981
 982
 983
 984
 985
 986
 987
 988
 989
 990
 991
 992
 993
 994
 995
 996
 997
 998
 999
 1000

1 The soluble lignin formed in the process of peroxide fractionation of larch wood is a light brown powder,
2 its yield reached 9-9.5 wt% on abs. dry wood. Elemental composition of lignin (wt%): C – 46.58, H – 3.79. The
3 simplest way of this lignin utilization is the obtaining of nanoporous materials.

4 The properties of enterosorbents obtained by alkaline treatment of peroxide lignins obtained from abies,
5 aspen and birch wood were studied early by authors (Garyntseva et al. 2011). The optimum method of
6 enterosorbents obtaining includes the treatment of peroxide lignins by the 0,4 % solution of NaHCO₃. The
7 adsorption activity of obtained enterosorbents (34.7-39.0 % for iodine, 88.4-89.2 mg/g for methylene blue and
8 183.0-212.0 mg/g for gelatin) is similar to adsorption activity of commercial enterosorbent “Polyphepan”. The yield
9 of enterosorbents is 73.4–76.1 % on the weight of initial lignin. Their characteristics correspond to pharmacological
10 standards. It can be expected that enterosorbents derived from larch wood lignin by the same way will possess by
11 similar properties.
12
13
14

15 For obtaining from lignin the nanoporous carbon the method of thermal activation can be used (Kuznetsov
16 et al. 2015). It was found that a partial gasification of wood and lignin in molten sodium and potassium hydroxides
17 allow to obtain the active carbons with developed porous structure. The textural characteristics of active carbons,
18 such as specific surface area, pore volume, macro-meso-micropores ratio, can be controlled by varying the nature of
19 the alkali and its content in the mixture (Kuznetsov et al. 2015).
20
21
22

23 The other possible way of lignin utilization is the preparation of organic and carbon aerogels (Aegerter et
24 al. 2011). They are highly porous materials combining a number of unique characteristics such as light weight, high
25 surface area and adjustable mesoporosity. Carbon aerogels may be used as high surface area adsorbents or catalyst
26 supports (Serp and Figueiredo 2009; Celzard et al. 2012; Szczurek et al. 2010; Job et al. 2006) and as electrodes for
27 double-layer capacitor (Amaral-Labat et al. 2012; Calvo et al. 2014).
28
29
30

31 Yearly it was suggested to use lignin for preparation of highly porous organic tannin-lignin aerogels
32 (Grishechko et al. 2013). So, the synthesis of new organic and carbon aerogels with the use of larch wood lignin is
33 the promising area of its potential application.
34

35 The obtained results in the field of larch biomass processing to valuable chemicals and functional organic
36 materials open new opportunities in green biorefinery of larch wood.
37
38
39

40 **Conclusion**

41 The new approach to green biorefinery of larch-wood is described, which allows to obtain a more wide
42 range of demanded products as compared to known methods of larch wood processing.
43

44 The suggested scheme of green biorefinery of larch-wood integrates the optimized methods of biomass
45 fractionation on DHQ, AG, MCC, soluble lignin and subsequent processing of polysaccharides to bioactive
46 polymers and lignin to nanoporous materials.
47
48
49

50 Described in the present paper the new methods of DHQ and AG extraction from larch-wood, peroxide
51 catalytic fractionation of extracted wood on MCC and soluble lignin along with early described original methods of
52 obtaining of bioactive sulfated polysaccharides and nanoporous materials from lignin use such non-toxic and less-
53 toxic reagents, as water, ethanol, hydrogen peroxide, acetic acid, sulfamic acid, urea.
54

55 The developed green method of one-step obtaining of DHQ and AG based on the water-ethanol extraction
56 of larch wood allow to obtain simultaneously the both compounds with high yields.
57

58 Based on kinetic studies and mathematical optimization of the process of extracted larch-wood peroxide
59 delignification in the presence of green catalyst TiO₂ the optimal conditions of wood effective fractionation on MCC
60 and soluble lignin were selected.
61
62
63
64
65

For green synthesis of bioactive functional polymers on the basis of sulfated MCC and AG we suggest to use non-dangerous sulfamic acid-urea mixture and methods of ion-exchange in water solutions.

Acknowledgements

The reported study was supported by Russian Science Foundation, grant N 16-13-10326.

This work is part of GDRI “Catalytic biomass transformation” between France and Russia.

References

- Aegerter MA, Leventis N, Koebel MM (2011) *Aerogels Handbook. Advances in sol-gel derived materials and technologies.* Springer-Verlag, New York
- Amaral-Labat G, Szczurek A, Fierro V, Stein N, Boulanger C, Pizzi A, Celzard A (2012) Pore structure and electrochemical performances of tannin-based carbon cryogels. *Biomass and Bioenergy* 39:274-282
- Anderson EM, Katahira R, Reed M, Resch MG, Karp EM, Beckham GT, Roman-Leshkov Yu (2016) Reductive catalytic of corn of corn stover lignin. *ACS Sustain Chem Eng* 4:6940–6950
- Babkin VA, Ostroukhova LA, D’jachkova SA, Svyatkin YuK, Babkin DV, Onuchin NA (1997) Waste-free complex processing of siberian and dahur larch biomass. *Chemistry for Sustainable Development* 45:105.
- Babkin VA, Ostroukhova LA, Malkov YuA (2000) RF Pat. 2158598
- Baryshnikov SV, Sharypov VI, Beregovtsova NG, Taran OP, Agabekov VE, Kuznetsov BN (2014) Thermal conversion of mechanically activated aspen-wood in sub- and supercritical ethanol medium. *J. Sib. Fed. University. Chemistry* 7:455-463
- Bessone M, Gallezot P, Pinel C (2014) Conversion of biomass into chemicals over metal catalysts. *Chem Rev* 114:827-1870
- Calvo EG, Lufrano F, Arenillas A, Brigandi A, Menendez JA, Staiti P (2014) Effect of unequal load of carbon xerogel in electrodes on the electrochemical performance of asymmetric supercapacitors. *Journal of Applied Electrochemistry* 44:481-489
- Celzard A, Fierro V, Amaral-Labat G (2012) Adsorption by Carbon Gels. In: Tascon JMD (ed) *Novel Carbon Adsorbents.* Elsevier, Oxford, pp 207-244
- Cherubini F, Jungmier G, Wellish M et al. (2009) Toward a common classification approach for biorefinery systems. *Biofuels, Bioproducts&Biorefining* 172:534-546
- Dagle VL, Smith C, Flake M, Albrecht KO, Gray MJ, Ramasamy KK, Dagle RA (2016) Integrated process for the catalytic conversion of biomass-derived syngas into transportation fuels. *Green Chem* 18:1880-1891
- Davis SC, Hay W, Pierce J (2014) *Biomass in the energy industry: an introduction.* BP p.l.c., London
- Ed de Long, Hidson A, Walsh P (2013) *Biorefinery. Task 42. Report International Energy Agency Bioenergy*
- Garyntseva NV, Sudakova IG, Kuznetsov BN (2011) Properties of enterosorbents obtained from acetic acid lignins of abies, aspen and birch wood. *J Sib Fed University Chemistry* 4:121-126
- Garyntseva NV, Sudakova IG, Kuznetsov BN (2015) Study of birch wood catalytic delignification by hydrogen peroxide at atmospheric pressure. *J Sib Fed University Chemistry* 8:422-429
- Gavrilov AB (2009) RF Pat. 2372095
- Gliozzi G, Innorta A, Mancini A, Bortolo R, Perego C, Ricci M, Cavani F (2014) Zr/P/O catalyst for the direct acid chemo-hydrolysis of non-pretreated microcrystalline cellulose and softwood sawdust. *Applied Catalysis B: Environmental* 145:24–33
- Goellner EM, Utermoehlen J, Kramer R, Classen B (2011) Structure of arabinogalactan from *Larix laricina* and its reactivity with antibodies directed against type-II-arabinogalactans. *Carbohydr Polym* 86: 1739-1744
- Grishechko LI, Amaral-Labat G, Szczurek A, Fierro V, Kuznetsov BN, Pizzi A, Celzard A (2013) New tannin-lignin aerogels. *Ind Crop Prod* 41:347-355
- Horhammer HS, Treasure TH, Gonzalez RW, Heiningen ARP (2014) Larch biorefinery: technical and economic evaluation. *Industrial and Engineering Chemistry Research* 53:1206–1213

- Hu F, Jung S, Radauskas A (2012) Pseudo-lignin formation and its impact on enzymatic hydrolysis. *Bioresource Technol* 117:7-12
- Hu L, Linc L, Wu Z, Zhou S, Liu S (2015) Chemocatalytic hydrolysis of cellulose into glucose over solid acid catalysts. *Applied Catalysis B: Environmental* 174:225–243
- Ishida K, Matsuda S, Watanabe M et al (2014) Hydrolysis of cellulose to produce glucose with solid acid catalysts in 1-butyl-3-methyl-imidazolium chloride ([bmIm][Cl]) with sequential water addition. *Biomass Conv Bioref* 4:323–331
- Isikgor FH, Becer CR (2015) Lignocellulosic biomass: a sustainable platform for production of bio-based chemicals and polymers. *Polym Chem* 6:4497-4559
- Job N, Heinrichs B, Lambert S, Pirard J, Colomer J, Vertruyen B et al. (2006) Carbon xerogels as catalyst supports: study of mass transfer. *AIChE J* 52:2663-2676
- Kislitsin AN, Mal'chikov EL (2009) RF Pat. 2346941
- Kuznetsov BN, Chesnokov NV, Ivanov IP, Veprikova EV, Ivanchenko NM (2015) Methods of porous materials obtaining from lignin and wood bark. *J Sib Fed University Chemistry* 8:232-255
- Kuznetsov BN, Chesnokov NV, Yatsenkova OV, Sharypov VI (2013) New methods of heterogeneous catalysis for lignocellulosic biomass conversion to chemicals. *Russian Chemical Bulletin* 62:1493-1502
- Kuznetsov BN, Kuznetsova SA, Danilov VG, Yatsenkova OV (2009) Influence of UV pretreatment on the abies wood catalytic delignification in the medium "acetic acid – hydrogen peroxide – TiO₂. *Reaction Kinetic and Catalysis Letters* 97:295-300
- Kuznetsov BN, Kuznetsova SA, Levdansky VA, Levdansky AV, Vasil'eva NYu, Chesnokov NV, Ivanchenko NM, Djakovitch L, Pinel C (2015) Optimized methods for obtaining cellulose and cellulose sulfates from birch wood. *Wood Science and Technology* 49:825-843
- Kuznetsov BN, Sharypov VI, Grishechko LI, Celzard A (2013) Integrated catalytic process for obtaining liquid fuels from renewable lignocellulosic biomass. *Kinetic and catalysis* 54:344-352
- Kuznetsov BN, Sudakova IG, Garyntseva NV, Djakovitch L, Pine C (2017) Kinetic studies and optimization of abies wood fractionation by hydrogen peroxide under mild conditions with TiO₂ catalyst. *React Kinet Mech Cat* 120:81-94
- Kuznetsov BN, Sudakova IG, Garyntseva NV, Djakovitch L, Pinel C (2013) Kinetic study of aspen-wood sawdust delignification by H₂O₂ with sulfuric acid catalyst under the mild conditions. *Reac Kinet Mech Cat* 110:271-280
- Kuznetsov BN, Taraban'ko VE, Kuznetsova SA (2008) New Catalytic Methods for Obtaining Cellulose and Other Chemical Products from Vegetable Biomass. *Kinetics and Catalysis* 49:517-526
- Kuznetsov BN, Vasilyeva NYu, Levdasky AV, Karacharov AA, Krylov AS, Mazurova EV, Bondarenko GN, Levdansky VA, Kazachenko AS (2016) Study by raman spectroscopy, xrd, sem, afm methods the sulfates of arabinogalactan obtained with the use of sulfamic acid. *Khimiya Rastitel'nogo Syr'ya (Chemistry of plant raw materials)* 4:43-48
- Kuznetsov BN, Vasilyeva NYu, Levdasky AV, Maximov AV, Kazachenko AS, Skvortsova GP, Djakovitch L, Pinel C (2016) Synthesis and study of copper-containing polymers of sulfated arabinogalactan. . *Khimiya Rastitel'nogo Syr'ya (Chemistry of plant raw materials)* 4:49-55
- Kuznetsova SA, Kuznetsov BN, Aleksandrova NB, Danilov VG, Zhizhaev AM (2005) Obtaining arabinogalactan, dihydroquercetin and microcrystalline cellulose using mechanochemical activation. *Chemistry for Sustainable Development* 13:261-268
- Kuznetsova SA, Yatsenkova OV, Danilov VG, Kuznetsov BN (2005) Oxidative delignification of larch wood in acetic acid-hydrogen peroxide-water medium in the presence of H₂MoO₄ catalyst. *Khimiya Rastitel'nogo syr'ya (Chemistry of Plant Raw Materials)* 4:35-39
- Li H, Fang Z, Smith Jr RL, Yang S (2016) Efficient valorization of biomass to biofuels with bifunctional solid catalytic materials. *Prog Energy Combust Sci* 55:98–194
- Liu D, Chen EY-X (2014) Integrated catalytic process for biomass conversion and upgrading to C₁₂ furoin and alkane fuel. *ACS Catal* 4:302–1310
- Liu Y, Chen L, Wang T, Zhang Q, Wang C, Yan J, Ma L (2015) One-pot catalytic conversion of raw lignocellulosic biomass into gasoline alkanes and chemicals over LiTaMoO₆ and Ru/C in aqueous phosphoric acid. *ACS Sustainable Chem Eng* 3:1745–1755

- 1 Ma C, Yang L, Wang W, Yang F, Zhao C, Zu Y (2012) Extraction of dihydroquercetin from *Larix gmelinii* with
2 ultrasound-assisted and microwave-assisted alternant digestion. *Int J Mol Sci* 13:8789–8804
- 3 Medvedeva EN, Babkin VA, Ostroukhova LA (2003) Arabinogalactan from larch—properties and
4 usage perspectives. *Khimiya Rastitel' nogo Syr'ya* (Chemistry of plant raw materials) 1:27-37
- 5 Naik SN, Goud VV, Rout PK, Dalai AK (2010) Production of first and second generation biofuels: a comprehensive
6 review. *Renewable and Sustainable Energy Reviews* 14:578-597
- 7 Pandey S (2015) Cellulases in conversion of lignocellulosic waste into second-generation biofuel. *International
8 Journal of Advanced Research* 3:392-399
- 9 Park S, Baker JO, Himmel ME, Parilla PA, Jonson DK (2010) Cellulose crystallinity index: measurement
10 techniques and their impact on integrating cellulose performance. *Biotechnol Biofuels* 3:10
- 11 Ramos LP (2003) The chemistry involved in the steam treatment of lignocellulosic materials. *Quim Nova* 26:863-
12 871
- 13 Ravber M, Knez Z, Skerget M (2015) Isolation of phenolic compounds from larch wood waste using pressurized hot
14 water: extraction, analysis and economic evaluation. *Cellulose* 22:3359–3375
- 15 Rowell RM (2012) *Handbook of wood chemistry and wood composites*, 2nd edn. CRC Press
- 16 Schutyser W, Van der Bosch S, Renders T, De Boe T, Koelewijn SF, Dewaele A, Ennaert T, Vikindereb O, Goderis
17 B, Courtin CM, Sels BF (2015) Influence of bio-based solvents on the catalytic reductive fractionation of
18 birch wood. *Green Chem* 17:5035-5045
- 19 Serp P, Figueiredo JL (2009) *Carbon Materials for Catalysis*. John Willey and Sons Inc, Hoboken
- 20 Serrano-Ruiz JC, Dumesic JA (2011) Catalytic routes for the conversion of biomass into liquid hydrocarbon
21 transportation fuels. *Energy Environ Sci* 4:83-99
- 22 Sharypov VI, Beregovtsova NG, Baryshnikov SV, Zhizhaev AM, Vosmerikov AV, Kuznetsov BN (2015) The
23 study of thermal conversion of mechanically activated aspen wood in the presence of high-silica zeolite
24 catalyst. *J. Sib. Fed. University. Chemistry* 8:190-201
- 25 Sixta H (2006) *Handbook of pulp*. Wiley-VCH Verlag GmbH & Co, Weinheim
- 26 Sjööstroöm E, Alern R (1999) *Analytical methods of wood chemistry. Pulping and papermaking*. Springer-Verlag,
27 Berlin
- 28 Sjöstrom E (1993) *Wood Chemistry- fundamentals and applications*, 2nd edn. Academic Press, New Work
- 29 Sudakova IG, Garyntseva NV, Yatsenkova OV, Kuznetsov BN (2013) Optimization of aspen wood delignification
30 by H₂O₂ with sulfuric acid catalyst. *J Sib Fed University Chemistry* 6:76-84
- 31 Szczurek A, Jurewicz K, Amaral-Labat G, Fierro V, Pizzi A, Celzard A (2010) Structure and electrochemical
32 capacitance of carbon cryogels derived from phenol–formaldehyde resins. *Carbon* 48:3874-3883
- 33 Tarabanko VE, Kaygorodov KL, Skiba EA, Tarabanko NV, Chelbina YuV, Baybakova OV, Kuznetsov BN,
34 Djakovitch L, Pinel C (2017) Processing pine wood into vanillin and glucose by sequential catalytic
35 oxidation and enzymatic hydrolysis. *J Wood Chem Technol* 37:43-51
- 36 Troshina AV, Roshchin VI (2014) Hydrocarbons of neutral substances from the branch bark of Siberian larch.
37 *Russian Chemical Bulletin* 63:2160–2168
- 38 Van den Bosch S, Schutyser W, Vanholme R, Drissen T, Koelewijn SF, Renders T, De Meester B, Huijgen WJJ,
39 Dehaen W, Courtin CM, Lagrain B, Boerjan W, Sels BF (2015) Reductive lignocellulose fractionation into
40 soluble lignin-derived phenolic monomers and dimers and processable carbohydrate pulps. *Energy Environ
41 Sci* 8:1748–1763
- 42 Vasilyeva NYu, Levday AV, Kazachenko AS, Skvortsova GP, Kuznetsov BN (2016) Modification of sulfated
43 arabinogalactan with amino acids by ion exchange method. *J Sib Fed University Chemistry* 9:20-28
- 44 Vasilyeva NYu, Levday AV, Kazachenko AS, Skvortsova GP, Kuznetsov BN, Djakovitch L, C. Pinel C (2016)
45 Synthesis of sulfated arabinogalactan derivatives with histidine and arginine *J Sib Fed University Chemistry*
46 9:318-325
- 47 Xiang LY, Mohammed MAP, Baharuddin AS (2016) Characterisation of microcrystalline cellulose from oil palm
48 fibres for food applications. *Carbohydrate Polymers* 148:11-20
- 49 Xu J, Xie X, Wang J, Jiang J (2016) Directional liquefaction coupling fractionation of lignocellulosic biomass for
50 platform chemicals. *Green Chem* 18:3124–3138

- Zhang X, Lei H, Zhu L, Qian M, Zhu X, Wu J, Chen S (2016) Enhancement of jet fuel range alkanes from co-feeding of lignocellulosic biomass with plastics via tandem catalytic conversions. *Appl Energ* 173:418–430
- Zhang X, Lei H, Zhu L, Wei Y, Liu Y, Yadavalli G, Yan D, Wu J, Chen S (2015) Production of renewable jet fuel range alkanes and aromatics via integrated catalytic processes of intact biomass. *Fuel* 160:375–385
- Zhang X, Lei H, Zhu L, Wu J, Chen S (2015) From lignocellulosic biomass to renewable cycloalkanes for jet fuels. *Green Chem* 17:4736–4747
- Zheng Y, Pan Z, Zhang R (2009) Overview of biomass pretreatment for cellulosic ethanol production. *Int J Agric Biol Eng* 3:51–68

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65