



Developments of the ecological tanning based on modified carbohydrates.

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Abstract

This work is based on the application development of biopolymers derived from modified starches, as per the study "Ecological tanning based on modified carbohydrates" presented at the III IULTCS EuroCongress Vicenza 2022, taking into consideration the tanning applied on hides deriving from liming processes of a different nature:

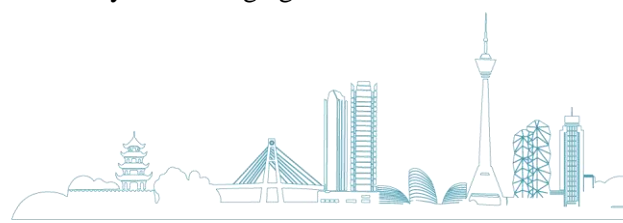
- traditional reductive liming with the use of sodium hydrosulphide and sodium sulphide
- reductive liming with the aid of thio-derivatives without the use of sodium hydrosulphide and sodium sulphide
- reductive liming as per the project "Odorless reductive liming, without thiols, thioglycolates and sodium sulphide"
- oxidative liming

In the previous study, all the application tests were carried out on hides derived from reductive liming based on thio-derivatives in the complete absence of hydrosulphides and sulphides, noting a good distribution and penetration of the tanning agent which improved performance when combined with various capping agents which included both organic substances (glutaraldehyde, oxazolidine and synthetic tannins) and inorganic substances (zeolites, silicates and non-metal free tanning agents). This work, on the other hand, will present the developments carried out on hides from different types of liming with the aid of new capping agents and different manufacturing procedures and will also be linked to the work "Odorless reductive liming, without thiols, thioglycolates and sodium sulphide", developed partly for starch tanning to improve the final quality of the leather.

Keywords: Oxidized Starch, Ecological Wet White, Natural based tanning

1. Introduction

Starch is the most abundant biopolymer in biosphere, produced by a large variety of plants. Due to its abundant, easy extraction method, low cost and renewable nature it became important not only for feeding but for all industrial sectors. Chemical modifications of starch, and other polysaccharides, are well studied and documented in the literature. Our goal is to produce and use a carbonyl rich oxidation product of starch as a tanning agent, in combination with other tanning agent, to obtain a leather with good merceological properties. The tanning effect of these type of agents is known in the literature and the reactivity towards the collagen is supposed to be like the classical aldehydes tanning agent with the





formation of an imine functionality that connect the protein structure of the hide with the polymeric structure of the oxidized starch [1-3].

There are two different sites, on the starch structure, where the reaction can take place in any glucose residue: C6 carbon atom and the C2-C3 site. The C6 carbon atom shows the typical reactivity of a primary alcohol toward oxidants species. The C2-C3 site, instead, shows the reactivity of a vicinal diol that can entails the scission of the carbon-carbon bond during the oxidation. The chemical functionalities that can be generated during the oxidation of starch are two: the carboxylic function and the partial oxidation product carbonylic function. Structure of the starch and of the possible oxidation products are showed in Fig.1, taking as example the linear structure of the amylose. The figure shows the limit structures of the possible oxidation products only for amylose, during the oxidation of the starch is also possible to have a mix of all functionality, that can be randomly located in the chain. The formation of an enolic group in C2-C3 in equilibrium with the ketonic tautomer is reported in literature but is not interesting for our purposes. In this process the origin, and therefore the

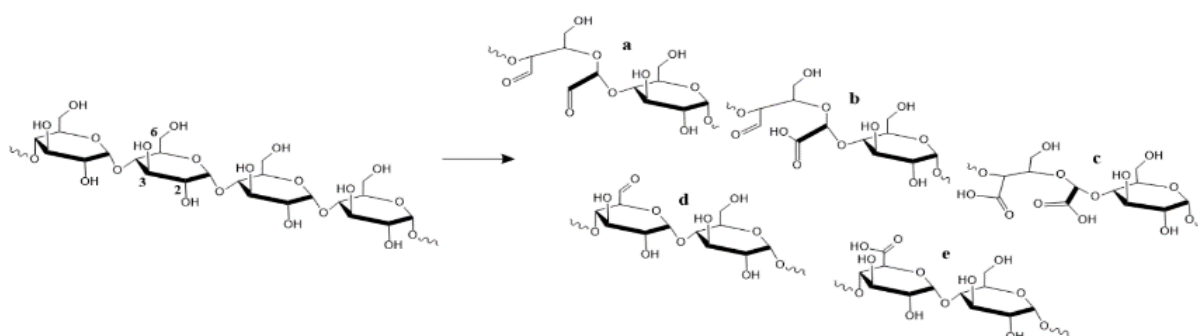


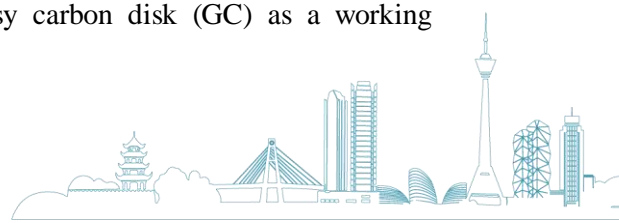
Figure 1. Possible oxidation products of the D-glucose residual in amylose, a) dialdehyde starch; b), c) product of partial and complete oxidation of C2-C3; d), e) carbonylic and carboxylic product of C6. Hemiacetals are also present.

nature, of the starch have a critical importance on the results. Oxidation happens on the surface of the starch granules dispersed in the liquid reaction medium and it is effective only in the amorphous regions and not in the crystalline one, so the effectiveness of the treatment depends on the amylose/amylopectin ratio, the crystallinity and the degree of polymerization of amylose [4,5].

The common ways to obtain the oxidation of starch are the polyols oxidation technique, that require the uses of an equivalent quantity of oxidants like sodium periodate (NaIO_4), hypochlorite, bromate, persulfate and hydrogen peroxide, TEMPO and other N-oxoammonium compounds. In this paper we put the attention on electrochemical methods for the oxidation of the starch that can be conducted with the use of green renewable energy sources. These methods do not require equivalent quantity of oxidant. Periodate is the laboratory scale classical way to obtain the carbonylic derivatives because it is pretty selective to the product known as dialdehydes starch (Fig. 1a) which is obtained through the oxidative ring opening on the C2-C3 functionality of the glucose residue [6,7]. The periodate is reduced to iodate, which is difficult to eliminate from the product and it is pollutant. The periodate has also the disadvantage of causing an extended depolymerization of the starch. The introduction of the electrochemical oxidation with the periodate opens the door to the use of catalytic amount of periodate thanks to the anodic regeneration of it [8,9]. Hypochlorite is more interesting for industrial purpose and cause an extensive oxidation that leads only the carboxylic products [10,11]. The method is also useful in electrosynthesis where the oxidating power of the hypochlorite ion, generated in situ at the electrode, can be mediated by the catalytic effect of the bromide ion [12] but the toxic gas Cl_2 is produced during the treatment.

2. Material and methods

2.1 Material: laboratory scale electrochemical tests are conducted in a 250 ml jacketed glass cell that can be equipped with a magnetic bar stirring system, a glassy carbon disk (GC) as a working





electrode, a Pt or graphite counter-electrode and a saturated calomel electrode (SCE) or Ag/AgCl like reference. In the preparative experiment agar gel septa was used at the scope to obtain a divided cell setup. Soluble potato starch, 2,2,6,6-Tetramethylpiperidine-1-oxyl (TEMPO), 4-acetamonium TEMPO (ACT), D-Glucose, 35% hydrogen peroxide, sodium carbonate, sodium phosphate, concentrated phosphoric acid, iron (III) ammonium sulphate dodecahydrate, disodium EDTA, sodium p-[(4,6-dichloro-1,3,5-triazinyl)amino]benzenesulphonate and solvents were purchased from Sigma Aldrich and used as received. Beamhouse products are supplied by Corichem.

2.2 Instruments: electrochemical test was performed by an AUTOLAB potentiostat/galvanostat equipped with a booster10A and Cyclic Voltammetry accessories. FT-IR spectroscopy was performed on a Nicolet iZ10 (Thermo Scientific) in ATR mode (SeZn crystal) or in transmission (KBr table). ^1H -NMR, ^{13}C -NMR and 2D-NMR correlation experiments was performed on a 600 MHz instrument (Bruker). HPLC analysis was performed in an Ultimate 3000 instrument (Thermos Scientific) equipped with a DAD and a 250x4.8 mm 5 μm 120 Å C18 HPLC column. Pilot scale experiments was performed on a divided-cells flow electrochemical pilot plant, the cathode act like the working electrode and is constituted by a graphite plate of the dimension of 0.04 m², the reservoir of the cell is constituted by a 10 L jacketed reactor. Counter electrode is an aluminium plate, the reservoir is a 10 L reactor. The two electrodes are divided by a nafion N324 selective membrane, the flow is guaranteed by two pumps equipped with flowstat. Tanning experiments were performed in a laboratory drum of 1.5 m of diameter.

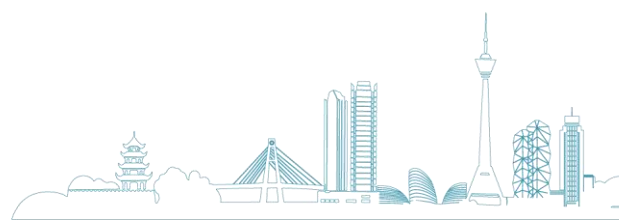
2.3 Determination of the electrochemical stability window of the substrate: a 20.6 g/L in water glucose dispersion in Na₂CO₃ 0.2007 M at 25 °C was introduced in the electrochemical cell and subjected to a 0-0.2V scan cyclic voltammetry at various scan rate.

2.4 Electrochemical preparations with TEMPO and derivates: cyclic voltammetry was conducted on a 1 mM TEMPO and Na₂CO₃ 0.2 M at different pH obtained by additions 1.5 M HCl with and without the presence of glucose (5.8 g/L) or starch (21.3 g/L) at 0.100 V/s and 25 °C. For quantitative preparation two GC and one Pt electrode were used. The reaction medium is constituted by a 50 ml of a carbonate/bicarbonate 0.2M buffer solution at pH=10.5 containing 39.1 mg of TEMPO, 10 g of starch and a 0.53 V vs SCE potential is applied for 15h. Cyclic voltammetries and preparative experiment with ACT were performed in the same condition substituting it to TEMPO in equal molarity. Products were characterized by FT-IR spectroscopy and by ^1H - ^{13}C COSY.

2.5 Electrochemical Fenton-Like oxidation with hydrogen peroxide: Cyclic voltammetries were conducted on a 0.1 M acetate or phosphate buffer at pH=3.0 containing 1 mM iron (III) ammonium sulfate solution and to the same solution with additional 1mM disodium EDTA at 25 °C scan rate 0.100 V/s. Then 2.0 ml of hydrogen peroxide was added and another cyclic voltammetry was recorded, then 501.4 mg of glucose was added and another voltammetry was recorded. The preparative test was conducted on 50 ml of buffer solution containing 47.23 mg of iron (III) ammonium sulphate, 86.24 mg of EDTA, 2 ml pf hydrogen peroxide and 401.45 mg of glucose was added, potential of -0.25 V was applied for 11220 s. The product was freeze-dried and ^1H -NMR was collected.

2.6 Pilot scale tests were conducted on the plant described above. The flow was maintained near 5.5 L/min for both the cells, different flow was applied to maintain the pressure throw the membrane equal for both sides, and we work in galvanostatic mode. The two tanks were filled each with 7 L of a 0.1M phosphate buffer obtained mixing sodium biphosphate with phosphoric acid to reach a determined pH and iron (III) ammonium sulphate dodecahydrate. The catholyte containing also 100g/L of soluble starch. The pH and quantity of iron salts are reported in Table 1. The circulation of the solution was turned on and the anolyte was added 76 g of 35% hydrogen peroxide. Then the galvanostat was turned on and set to allow to pass 4 A of current. Periodically, a sample was collected to determine the aldehyde and hydrogen peroxide content.

2.7 Tanning experiments were performed in a laboratory drum. Pelt was delimited and bated with common standard procedures. Samples were taken to perform analysis.





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Quantity [%]	Product	Temp. [°C]	Time [min]	Notes
20	Water	20		
0.025	Sulfonate fatliquor		20	
10	2.6d (tanning agent)		30	
10	2.6d (tanning agent)		Overnight	pH=4.2
0.5-1	Formic acid		30-60	If required, depending from the capping agent to obtain pH=3.6
	Capping agent			
0.5-2	Sodium bicarbonate			If acid was required; quantity and time for obtain the target pH of 4.7
				Drain
100	Water	20		
0.1	EDTA		20	Drain
Splitting, the rest of the process was conducted only on grain				
100	Water	35		
8	Filler		40	
4	Melaminic resin		30	
1.5	Neutralizing tannin		30	
4	2.6d (tanning agent)		30	
14	Mixed tannin and syntan		40	
5	Mixed Fatliquor		60	
200	Water	50		
1	Formic acid		Overnight	
2	Charge inverter		60	Drain and wash
50	Water	30		
14	Mixed tannin and syntan		60	
12	Mixed fatliquor		40	
0.5	Ammonia		5	
2.5	Dye staff		60	
4	Melamin resin		40	
200	Water	50	15	
2	Formic acid		40	Drain and wash

Table 1. Tanning procedure for the test 2.7 a, b, c, d1, d2, d3, d4.

For the tests we used pelt coming from a reductive liming process with the aid of thio-derivatives without the use of sodium hydrosulphide and sodium sulphide and the following capping: no capping (2.7a), 3% of sodium p-[(4,6-dichloro-1,3,5-triazinyl)amino]benzensulphonate (test 2.7b), 1.5% of glyoxal (test 2.7c). The 2.7d test series was conducted using a 4% of a silicate derivatives as a capping agent and starting from pelts obtained from liming: the same as the previous ones (2.7d1), traditional reductive (2.7d2), odorless reductive liming without thiols, thioglycolates and sodium sulphide (2.7d3), oxidative liming (2.7d4). Test 2.7e corresponds to a performance target obtained by conventional glutaraldehyde tanning process.

2.8 Chemical oxidation tests were conducted preparing a solution of soluble starch, iron salt and a buffer, for example phosphate or acetate, in order to obtain a pH from 3 to 6. Iron salt and Hydrogen peroxide is then added on top of that in small portions during several hours to maintain the temperature under 50°C, then is left to stir for 8 h. The product obtained was tested in tanning like 2.6d.





2.9 Determination of the aldehyde content in oxidized carbohydrate (CC, Carbonyl concentration) was conducted via titration according with [13]. 5 ml of sample was introduced into a 250 ml becher, on top of that it was added 50 ml of distilled water and the pH was adjusted to 3.2 with 0.1 M HCl and is heated to 40 °C in a thermostatic bath. Therefore 60 ml of a 0.36 M Hydroxylamine Hydrochloride and 0.1 M NaOH solution is added, the becher was covered with aluminium foil and allow to stay a 40 °C for 4h. Than the solution was pH-metrically titred to pH=3.2 with a 0.1 HCl solution. The carbonyl content, as percentage of C-OH groups converted to carbonyl groups, is calculated as sequent: $CC(\%) = \frac{(V_B - V_S)}{0.01851 \times m}$ where m is the dry weight of the sample contained in the volume titrated. Tensile strength and extension, tear propagation force and determination of distension and strength of the leather grain were performed respectively according to methods ISO 3376 [A], ISO 3377-2 [A] and ISO 3379 [A] by an external certified laboratory.

3. Results and Discussion

Dextrose was selected like model substrate to conduct the preliminary screening experiments. All the electrochemical processes that we use are electro-catalyzed, that mean that the electron transfers involved in the oxidation of the various carbohydrates used are conducted or mediated by species that are regenerated at the electrode, so the oxidation of the starting materials do not happen at the electrode-solution interphases but in the bulk region. The first electrocatalytic process investigate was the N-oxoammonium compounds using TEMPO and ACT. Oxoammonium compounds are very selective to the oxidation of alcohols to aldehydes, avoiding overoxidation to carboxylic groups. Cyclic voltammtries at various scan rate of the sole substrate, in Fig. 2a, reveal no redox processes affecting glucose in that window.

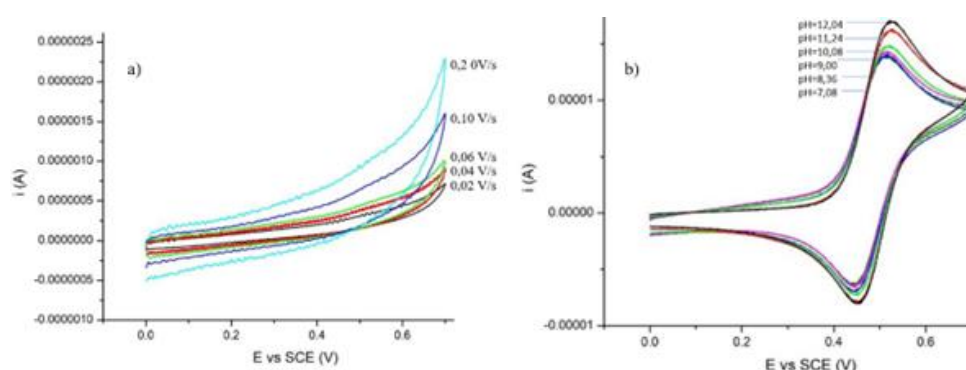
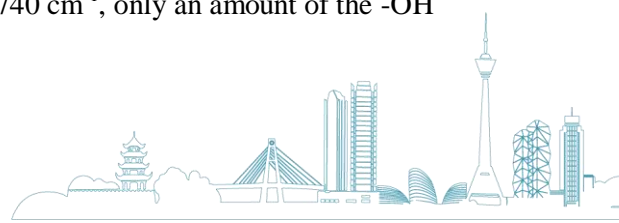


Figure 2 Cyclic voltammtries of various preliminary tests recorded according to the procedure a) 2.3; b) 2.4[14]

Subsequently the optimum pH for the catalyst is determined via cyclic voltammetry on a TEMPO solution, the analysis reveal a reversible redox process with an increasing of the anodic and cathodic peak at the increasing of the pH (Fig. 2b), same results were obtained for ACT. The addition to glucose to the system, Fig. 2c, cause a sensible increasing of the anodic current and the disappear of the cathodic peak. The introduction of a reducing species that react with the oxidized catalyst, but is not involved in a redox process at the electrode in the studied potential window, make the process clearly irreversible because the reduction of TEMPO is conducted by glucose and not by the electrode. Is also possible to observe an increasing of the peak current, because of the concentration of the reduced catalyst in the interphase is now increased by the reducing action of glucose. More concentration means more electron transfer and consequently more current, obtaining a cyclic voltammetry that clearly shows that we have an electrocatalytic process. The preparative test is designed according to the information obtained by the previous experiments. Results in Fig.4 show that the TEMPO acts like a better catalyst because in 1000 s it allows to pass 149.21 C against ACT that only passes 98.01 C, so it need more time to obtain the same results. Infrared analysis (Fig.4c) reveals the formation of aldehydic group because of the peak at 1740 cm^{-1} , only an amount of the -OH





groups involved in the oxidation process are oxidized to carbonyl group, another part remains unreacted and another portion is overoxidized to carboxylate, identified by the peak at 1636 cm^{-1} . NMR experiment confirms these results and shows the presence of carboxylic hydrogen revealed by the signal at 7.93 ppm on the F1 channel in Fig.4d.

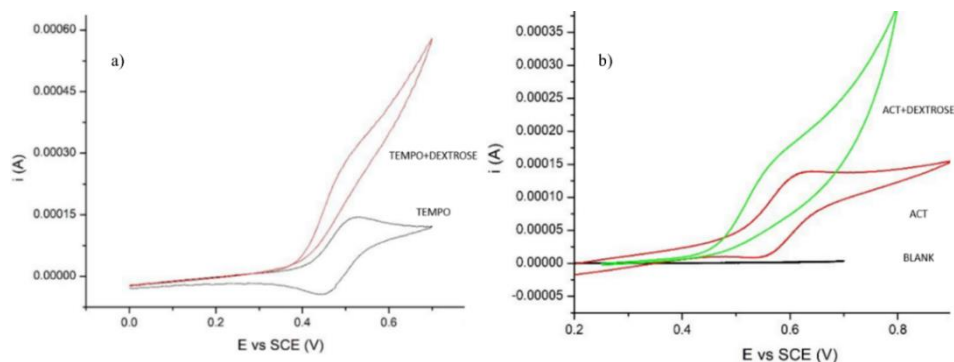


Figure 3. Cyclic voltammeteries of TEMPO and ACT catalysed electrolysis of dextrose.

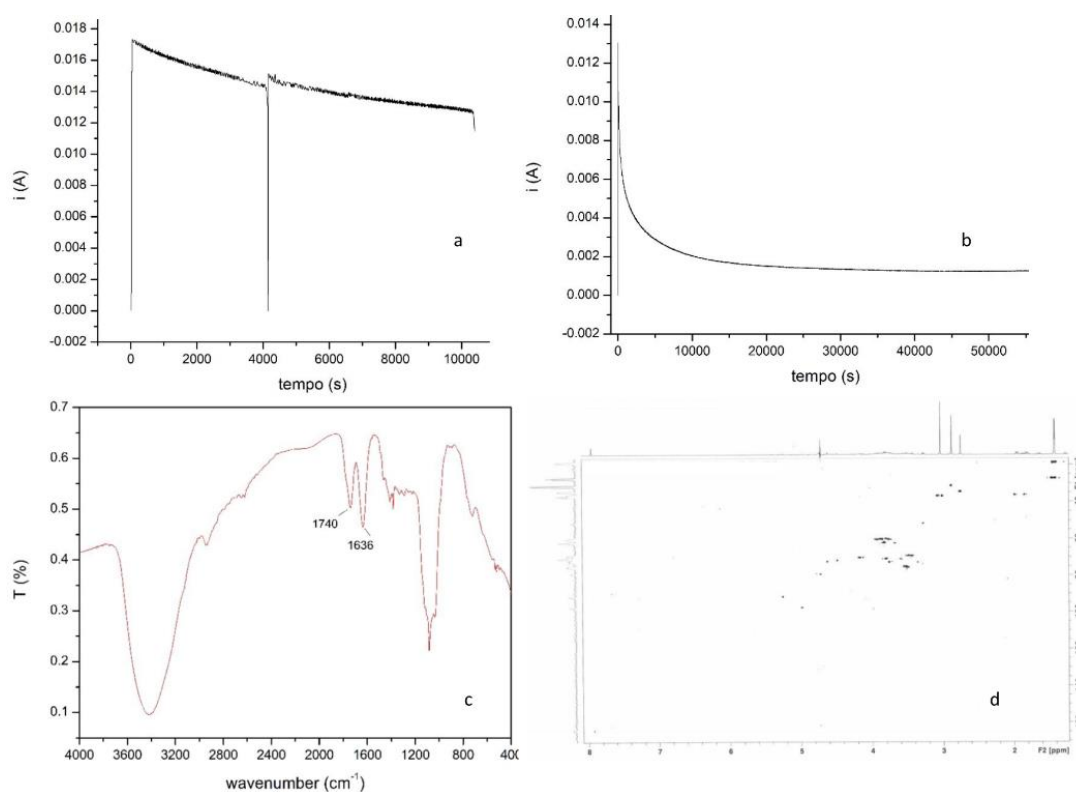


Figure 4. Graphix of current versus time of the preparative tests of oxidation of dextrose with a) TEMPO, spike is caused by operator error; b) ACT. c) and b) show respectively the FT-IR ATR and ^1H - ^{13}C COSY experiments on the TEMPO oxidized freeze-dried products.

The possibility of use a much strong oxidating method to obtain the oxidation of starch was also investigated, an oxidating method called electro-Fenton reaction, a technique based on Fenton chemistry. Simplifying the mechanism of the conventional Fenton reaction Fe (II) is oxidized by hydrogen peroxide to Fe (III) producing the hydroxide radical, a strong oxidating species, and a hydroxide ion, therefore the Fe (III) is reduced back by hydrogen peroxide to Fe (II) producing hydroperoxide radical and an hydronium ion, restarting the catalytic cycle. That means that a quantity of hydrogen peroxide is consumed to restore Fe (II) in a slow step that produce a lesser oxidizing species. In addition, other reactions which require the presence of Fe (III) that are not described here, contribute to depress the reactivity of Fenton oxidation [15]. The idea is to continuously reduce the Fe





(III) at the electrode in order to maintain its concentration as low as possible to prevent undesired reactions, promote the reactivity and, eventually, electrogenerated the hydrogen peroxide. Because the reduction of Fe (III) in this system is not performed by hydrogen peroxide there is not the production of H^+ ion that neutralizes the OH^- ion produced by the Fe (II) oxidation, so there is the necessity of a buffer to maintain stable the pH. We conduce cyclic voltammetries on a Fe (III) solution with and without the presence of disodium EDTA. As we see in Fig. 5a the presence of a complexing agent makes the process more reversible and it is possible that the complexation makes the Fe (II) a more powerful reducing agent because of the shift of the cathodic peak to more negative potential.

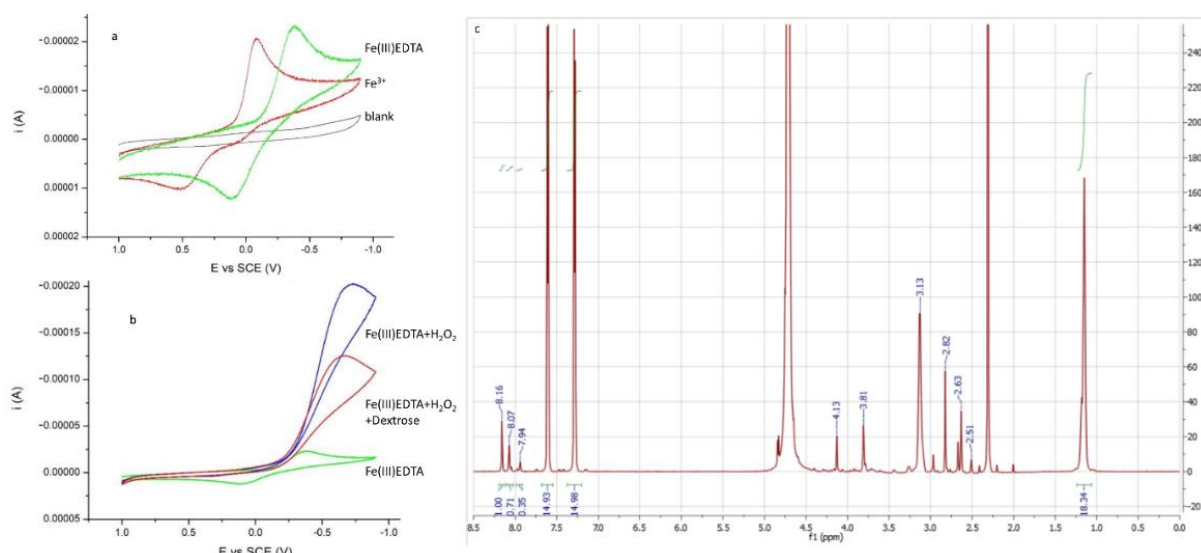


Figure 5 cyclic voltammetries and 1H -NMR (D_2O , 400 MHz) obtained from the procedures 2.5 [14]

A more reversible shape is obtained substituting the acetate buffer, used to collect the voltammogram Fig. 5a, with phosphate buffer, in Fig. 5b, because the acetate interferes in the complexation of iron with EDTA acting like a competitive ligand. The addition of hydrogen peroxide changes the cyclic voltammetry from a reversible shape to the irreversible one shown in Fig. 5b because of the Fe (II) is not oxidized back at the electrode anymore but by hydrogen peroxide, increasing the Fe (III) concentration in the interphase and consequently increasing the peak current. The addition of dextrose drops the cathodic peak current because of the hydroxy radical now react with the carbohydrates and do not involve in the possible side reaction of oxidation of Fe (II) to Fe (III), causing the decreasing of the concentration of the last one at the electrode; this prove that dextrose is involved in the process.

Analyzing the voltammetries, an applied potential of $E_{app} = -0.250$ V was selected for preparative scale experiment in order to obtain the reduction of iron but not the hydrogen peroxide one. In the test 145.4 C were passed to the cell with an initial current of 21.0 mA that drop to 6.3 mA near to the end of the experiment. The 1H -NMR of the product of the test permits to qualitative identify the aldehyde groups formed during the test that correspond to the signals at chemical shift of $\delta = 7.94, 8.07, 8.16$ ppm. This result permits to pass with this setup to the pilot scale experiment.

The pilot scale test was performed in the pilot plant showed in Figure 6. The results of the various experiments are showed in Table 2. As we can see pH and catalyst content are determinant, less acid pH increases the Fenton reactivity but it is not possible to increase the pH a lot, in order to avoid

The pilot	pH	Iron salt	Aldehyde
2.6a	3.2	6.3	4.5
2.6b	3.2	20	8.19
2.6c	4	6.3	5.67
2.6d	4	20	9.45

Table 2. pH, iron salt and aldehyde content of the final product of the four tests performed on the pilot scale plant.





hydrogen peroxide self-degradation and iron compounds precipitation. The optimum pH range is determined to be between 3.2 and 4 by the use of speciation diagrams of iron in water. With the increasing of the catalyst concentration there is an increment of the aldehyde content. We cannot abuse in its quantity because is difficult to remove from the final product and imparts a red colour that is transferred to the hide during tanning and can increase the presence of iron defect. The better results are obtained with the test 2.6d that is repeated 10 times for producing materials for the subsequent tanning tests and for proof the reproducibility of the preparation. The classical Fenton chemistry, used in experiment 2.8, gives quantity of carbonyl group of 5.42 % or higher obtained in our tests. However, this type of reaction appears to be difficult to control due to the exothermic character, with overoxidation to carboxyl. Fig. 7 puts in evidence the different ratio between the carbonyl (1740 cm^{-1}) and carboxyl (1712 cm^{-1}) signals. It is possible to see the high degree of oxidation produced by classical Fenton approach where the carboxyl signal is higher than carbonyl one, that means the problem is overoxidation.

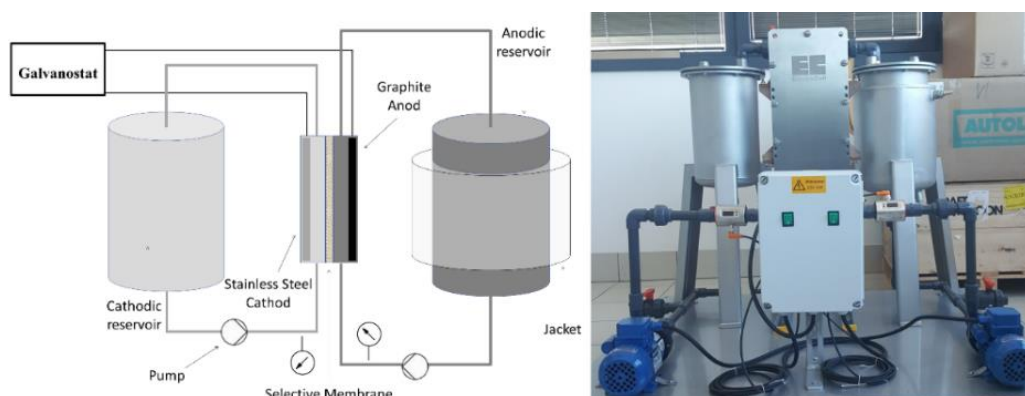


Figure 6 Schematic representation and picture of the pilot scale plant.

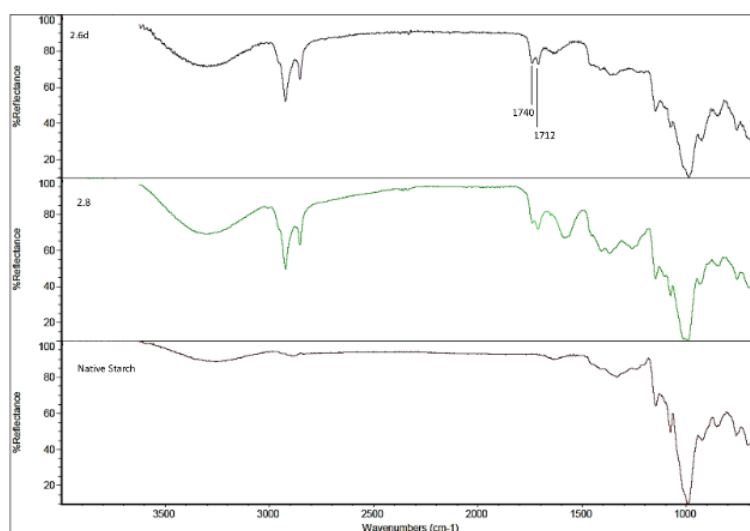


Figure 7. Comparison between the normalized FT-IR spectrum of 2.8, 2.7d and native starch.

Obtained results in term of leather quality are reported in Table 3 and 4. It is noted, by comparison between the Tg of test 2.7a and tests 2.7b,c that the glyoxal and the triazine derivative do not act as good capping agents, depressing the tanning effect 2.6d. The best results are obtained by using silicate as a capping agent when the tanning process takes place on pelts from liming processes without sulphides (test 2.7d1) and odorless (test 2.7d3). The chemical-physical properties of the leather obtained from these two tests are in line with those of the standard article 2.7e taken as a target. Also, in terms of merchandise evaluation, the leather obtained from tests 2.7d1 and 3 are satisfactory.





Sample	T _g after tanning [°C]	T _g crust [°C]	Tear propagation force [N]	Tensile Test		Distension and Strength			
				Tensile Strength [N/mm ²]	Elongation at max. force [%]	Grain distension [mm]	Load at grain crack [N]	Distension at crack [mm]	Load at crack [N]
2.7a	65	75	87.0	9.64	40.9	7.8	221	9.0	274
2.7b	62	65	82.1	8.51	38.2	7.5	201	8.8	250
2.7c	63	65	80.9	8.82	37.3	7.7	210	8.9	260
2.7d1	71	81	127.0	16.23	53.2	8.8	341	11.1	532
2.7d2	65	73	110.3	15.27	48.3	8.8	270	9.9	485
2.7d3	69	80	125.8	16.54	50.3	9.0	300	10.3	510
2.7d4	62	67	100.5	14.05	45.3	8.8	240	9.5	460
2.7e	75	81	108.3	15.11	64.0	9.4	356	11.7	488

Table 3. Chemical-physical property of obtained leathers.

Sample	Merchandise evaluation (from 1 to 10)			
	Softening	Firmness	Roundness	Tinctorial yield
2.7a	5.3	8.3	8.0	5.6
2.7b	5.0	6.0	6.5	7.0
2.7c	6.0	6.0	6.5	7.0
2.7d1	5.7	5.7	5.9	7.3
2.7d2	5.5	6.0	5.0	6.0
2.7d3	7.5	7.0	8.0	7.3
2.7d4	8.0	6.0	8.0	7.0
2.7e	8.2	5.2	8.2	5.8

Table 4 Merchandise evaluation of obtained leathers.

4. Conclusion

In conclusion this work proves the possibility of obtaining oxidized starch by the use of electro-Fenton approach that gives better results, in terms of carbonyl content, than classical Fenton or electrocatalytic TEMPO oxidation. The method does not produce toxic species and do not need further purification like chlorine or periodate methods. The product of the oxidation has proven to be a useful tanning agent if it is applied with an appropriate tanning procedure that involve the use of capping agent. In this sense the better agent is proven to be the silicate that provide a leather that possess chemical-physical properties very similar to standard glutaraldehyde tanned leather.

5. Acknowledgement

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