Hydroxamic cellulose fibre as a chelating cation exchanger

Grdinić, V.; Kujundžić, N.

Source / Izvornik: Croatica Chemica Acta, 1981, 54, 109 - 113

Journal article, Published version Rad u časopisu, Objavljena verzija rada (izdavačev PDF)

Permanent link / Trajna poveznica: https://urn.nsk.hr/um:nbn:hr:163:725289

Rights / Prava: In copyright/Zaštićeno autorskim pravom.

Download date / Datum preuzimanja: 2025-04-03



Repository / Repozitorij:

Repository of Faculty of Pharmacy and Biochemistry University of Zagreb



CCA-1265

YU ISSN 0011-1643 UDC 547.458.81:543.854.748 Original Scientific Paper

Hydroxamic Cellulose Fibre as a Chelating Cation Exchanger

V. Grdinić and N. Kujundžić

Department of Chemistry, Faculty of Pharmacy and Biochemistry, University of Zagreb, 41000 Zagreb, Croatia, Yugoslavia

Received June 25, 1980

Hydroxamic cellulose fibre with a high content of hydroxamic groups was prepared. The extent of hydroxamation was established by determination of nitrogen. The sorption capacity of the fibre was estimated to be 2.1 mekv/g. The hydroxamic cellulose fibre represents a selective sorbent for fast preconcentration of traces of ions. The effect of the mass of the sorbent, salt effect and contact time on sorption, were studied by using 5—30 mg of the hydroxamic cellulose in 25 ml from 1 to 10 ppm iron(III) solutions. Preliminary results show that hydroxamic cellulose may be used in environmental analytics.

INTRODUCTION

Solid chelating ionic exchangers with cellulose as a base have often been used recently for preconcentration and determination of traces of ions in environmental analytics. A number of compounds with different chelating groups such as: salicylic $\operatorname{acid^{1,2}}$, (β) -alanine-N,N-diacetic $\operatorname{acid^3}$, 8-hydroxyquinoline⁴, 1-(2-hydroxyphenylazo)-2-naphthol^{5,6}, 1,2-dihydroxybenzene-3,5-disulphonic $\operatorname{acid^7}$, o-aminophenol, 1-(2-hydroxyphenylazo)-2-naphthol, 1-(2-hydroxyphenylazo)-2-naphthylamine-N,N-di acetic $\operatorname{acid^8}$, chromotropic $\operatorname{acid^9}$, NN-ethylenedi-(-o-hydroxyphenyl)glycine, quinalizarin, arsenazo III, 1-(2-hydroxyphenylazo)-2,4-phenylenediamine-N,N,N'N'-tetraacetic $\operatorname{acid^{10}}$, and iminodiacetic $\operatorname{acid^{11,12}}$ were prepared and examined.

The synthesis of a poly(hydroxamic acid) ion exchange resin from cross-linked poly(acrylonitrile) is described 13 . A hydroxamized cellulose powder with a low content of hydroxamic groups was prepared from a starting material of carboxy-cellulose containing about 6 wt. 0 / 0 0 of carboxyl groups 14 .

In this paper we wish to report the preparation and some analytical properties of chelating cellulose with a high content of hydroxamic acid as a functional group.

EXPERIMENTAL

Chemicals

Oxidized regenerated cellulose, Surgical (Johnson Johnson, New Yersey, USA) with about 18 wt. 0 / 0 0 carboxy groups was used as a starting material for the preparation of hydroxamized cellulose. Hydroxamized cellulose fibre was prepared as subsequently described. Hydroxylamine hydrochloride and methanol were of Merck p. a. grade. All other chemicals were of analytical grade and were used without further purification. The water used in all experiments was redistilled from aqueous acidified potassium permanganate solution.

Methods

The structures of the products were established by elemental analysis and spectral data. Infrared spectra were recorded on a Perkin-Elmer spectrophotometer Model 457 using KBr pellets. Water content was determined gravimetrically after the samples had been dried 20 hours at room temperature in a vacuum thermostat. Sorption of iron(III) ions was followed spectrophotometrically at 480 nm with a Zeiss-Opton spectrophotometer PMQ II using ammonium thiocyanate as a reagent 15 and 1.000 cm quartz cells.

Preparation of hydroxamized cellulose

Hydroxylamine hydrochloride (3.63 g) was dissolved in 40 ml of cooled anhydrous methanol and neutralized with an equimolar quantity of sodium methylate (2.81 g) dissolved in 50 ml of methanol.

The mixutre was cooled for 2 hours in a refrigerator and precipitated sodium chloride was filtered off.

Surgical cellulose fibre (Celuronic acid) (1.3 g) was suspended in 20 ml of methanol cooled down to $4\,^{\circ}\text{C}$ and added slowly to the cold hydroxylamine solution. The mixture was stirred for 5 hours at room temperature and then allowed to stand in a refrigerator overnight.

The fibreous mass was decanted, washed with 20 ml methanole, filtered off and washed 5 times with 20 ml methanol. The product was dried in air and stored in vacuum dessicator. The water content was usually $3.5^{\circ}/_{\circ}$. Characteristic IR bands (KBr): 330—3600 (OH), 2900 (CH, CH₂), 2710 (NH), 1735 (C=O), 1550—1700 (C=O, NH), 1420 (CH₂), 1290, 1240 and 800 cm⁻¹.

N content: 2.98%

RESULTS AND DISCUSSION

Sorption capacity

The IR spectra, nitrogen content and colored reaction of iron(III) ions with the product indicate that hydroxamic groups were incorporated into the anhydroglucose unit of the cellulose molecule. The yield is 12.6 wt.% CONHOH. According to the nitrogen content the sorption capacity may be estimated as 2.13 mekv/g of the dried sorbent.

Solubility of the fibres

The fibres of the sorbens are not soluble in neutral and acidic medium. It is soluble in alkaline solutions and subsequent hydrolysis of the hydro-xamized cellulose was observed. Further experiments are in progress to throw more light on the events that take place in this system after hydroxamized cellulose is dissolved in alkaline solution.

Limit of identification an selectivity for somes ions

Some of metal ions with hydroxamic cellulose forming colored products¹⁶. Hydroxamic cellulose may be used as an analytical reagent and/or as a sorbent.

For such ions the identification limits were estimated by the »resin spot test« method¹¹ applying semistatistical analysis¹8. The obtained values for identification limits are: 5.3×10^{-7} g Au³+, 1.3×10^{-6} g Cu²+, and 8.0×10^{-7} g Fe³+. The colour of the products were different: red-brown (Fe³+), yellow-green (Cu²+) and gray (Au³+). The following ions do not give visible products with hydro-xamized cellulose at a concentration of 10 µg/0.05 ml: Ag⁺, Al³+, Bi³+, Cd²+, Cr³+, Hg²+, La³+, Mn³+, Mn²+, Ni²+, Pb²+, Sb³+, Th⁴+, Ti³+, UO₂²+ and Zr⁴+.

Salt effect

Effect of different salts on the sorption of iron(III) ions on hydroxamic cellulose is shown in Table I. It is clear that some of the salts such as $CdCl_2$, KCl, $LaCl_3$, $Th(NO_3)_4$, $UO_2(NO_3)_2$, and $ZrOCl_2$ increase the sorption of iron(III) ions on the hydroxamic cellulose. $AgNO_3$ exibits no effect, however $Al_2(SO_4)_3$ showed a small decreasing effect. In the presence of $Cu(NO_3)_2$ the sorption of iron(III) ions is lacking probably because of strong sorption of copper(II) ions.

Salt		mg of the	iron(III)*
Formula	mg in 25 ml	sorbent in 25 ml	sorbed µg/ml
_		20.0	3.39
$AgNO_3$	20.0	21.2	3.39
$Al_2(SO_4)_3$	20.5	20.5	3.11
CdCl ₂	20.0	20.4	4.73
Cu(NO ₃) ₂	20.0	21.0	0.00
KCl	21.5	20.5	4.22
LaCl ₃	20.8	20.5	4.73
Th(NO ₃) ₄	21.2	20.1	4.69
$UO_2(NO_3)_2$	20.5	20.0	3.90
$ZrOCl_2$	21.0	20.8	4.10

^{*} In all experiments the starting concentration of iron(III) was 5.95 $\mu g/ml$

Sorption characteristics of the hydroxam cellulose

Hydroxamic cellulose is a hydrophilic very porous fibre like chelating ionic exchanger. Our qualitative tests show that it is a selective sorbent with special affinit for preconcentration of iron(III) ions. Sorption of iron(III) ions from acidic solution was a very fast process as is shown in Figure 1. In this experiment iron(III) ion concentration was 5.86 ppm with 5.0—5.2 mg of the sorbent. The sorption is affected by the quantity of sorbent quantity as is shown in Figure 2. The larger amount of the sorbent in same shaking time sorbed larger amount of iron(III)-ions. The optimal conditions for the sorption of iron(III) ions on hydroxam cellulose are the subject of a study which is in progres in our laboratory.

Acknowledgement. — The authors are grateful to the Council for Research of Croatia (SIZ II) for support and to the Alexander von Humboldt Foundation for the Opton PMQ II spectrophotometer.

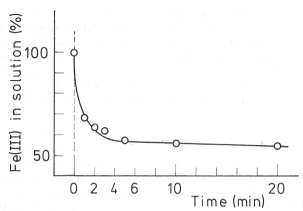


Figure 1. The sorption of iron(III)-ions on hydroxamic cellulose as a function of time. Starting concentration of iron(III)-ions was 5.86 µg/ml. Mass of the sorbent was 5.0-5.2 mg in 25 ml.

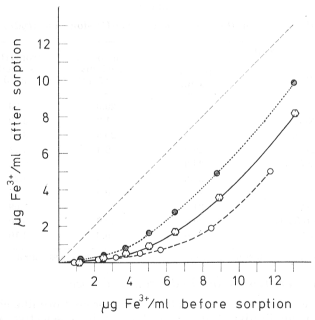


Figure 2. Effects of the mass of the sorbent on the sorption of iron(III)-ions on hydroxamic cellulose. $\square=30$ 0—31.0 mg of the sorbent; $\bigcirc=20.0$ —21.6 mg of the sorbent; $\bigcirc=9.8$ —12.0 mg of the sorbent. Shaking time was 10 min.

REFERENCES

- P. Burba and K. H. Lieser, Z. Anal. Chem. 279 (1976) 17.
 K. H. Lieser, M. Forster, and P. Burba, Z. Anal. Chem. 284 (1977) 199.
 I. Kojdl, J. Pract. Chem. 38 (1969) 851.

- H. Burba and K. H. Lieser, Angew. Macromol. Chem. 50 (1976) 151.
 P. Burba and K. H. Lieser, Z. Anal. Chem. 286 (1977) 191.
 F. Ambe, P. Burba, and K. H. Lieser, Z. Anal. Chem. 295 (1979) 13.
 P. Burba, M. Griesbach, and K. H. Lieser, Z. Anal. Chem. 248 (1977)
- 8. P. Burba and K. H. Lieser, Angew. Macromol. Chem. 64 (1977) 197.

- 9. K. H. Lieser, H. M. Rober, and P. Burba, Z. Anal. Chem. 284 (1977) 361.
- 10. P. Burba, M. Rober, and K. H. Lieser, Angew. Macromol. Chem. 66 (1978) 131.
- 11. Zs. Horvath, K. Falb, and M. Varju, At. Absorption Newslett. 16 (1977) 152.
- 12. Zs. Horvath and Gy. Nagydiosi, J. Inorg. Nucl. Chem. 37 (1975) 767.
- 13. F. Vernon and H. Eccles, Anal. Chim. Acta 82 (1976) 369.

- 14. K. Kotsuji and S. Hayashi, Japan Analyst. 23 (1974) 802.
 15. J. Bognar and M. P. Szabo, Microchim. Acta (1968) 957.
 16. E. Gagliardi and H. Raber, Mh. Chem. 93 (1962) 360.
 17. M. Fujimoto, Bull. Chem. Soc. Japan 27 (1954) 48.
- 18. V. Grdinić and G. Špoljarić, Farm. Glas. 36 (1980) 41.

SAŽETAK

Vlakna hidroksam celuloze kao novi kelirajući kationski izmjenjivač

V. Grdinić i N. Kujundžić

Pripremljena su vlakna hidroksam celuloze iz karbokarboksi celuloze s visokim sadržajem hidroksamskih skupina. Određivanjem sadržaja dušika ustanovljen je postotak hidroksamacije i procijenjen sorpcijski kapacitet vlakna koji iznosi 2,1 mekv/g. Vlakna hidroksam celuloze predstavljaju selektivni sorbens za brzo prekoncentriranje tragova iona. Efekt mase sorbensa, stranih soli i trajanja kontaktiranja na sakupljanje je ispitan primjenom 5—30 mg sorbensa u 25 ml otopine 1—10 ppm željeza(III). Preliminarna ispitivanja pokazuju potencijalnu mogućnost primjene hidroksam celuloze u analitici prirodnih sustava.

ZAVOD ZA KEMIJU FARMACEUTSKO-BIOKEMIJSKI FAKULTET SVEUČILIŠTA U ZAGREBU

Prispjelo 25. lipnja 1980.