## Cellulose: chemical modification

**CHEM-E2140** 



Eero Kontturi 30<sup>th</sup> September 2021

## Learning objectives

#### After this lecture, you will be able to:

- Distinguish the common pathways of cellulose modification: esterification and etherification
- List the most common cellulose derivatives, how they are made, how they dissolve, and what they are used for
- Be aware of the concept of regioselectivity
- Calculate the degree of substitution



## **Outline**

- (1) Chemical modification of cellulose motivation
- (2) Background: terminology, challenges
- (3) Degree of substitution: how to calculate
- (4) Esterification of cellulose
- (5) Etherification of cellulose
- (6) Regioselectivity in chemical modification of cellulose



# Motivation for cellulose modification

- Preparation of substances that have different properties from cellulose, yet they are of renewable origin and (to an extent) biodegradable
  - One of the most important properties is that most cellulose esters and ethers are thermoplastic (cellulose is not)
- Modified cellulose, i.e., cellulose derivatives often possess properties that are not easily achieved with totally synthetic polymers
- (With nanocellulose) modify the surface of nanocelluose to achieve better compatibility with its environment (composites etc.) or induce various functionalities (responsivity etc.)



## **Basic concepts**

• The idea of chemical modification of cellulose is to introduce functional groups in the cellulose backbone

 Usually achieved by substituting the protons in the hydroxyl groups of cellulose to a varying extent



## **Basic concepts**

#### (1) Homogeneous modification

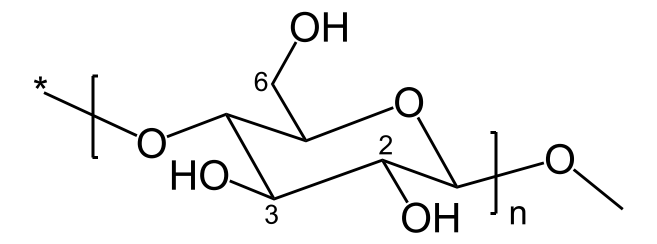
- Cellulose is dissolved and individualized cellulose chains are modified in a homogeneous solution
- → Uniform, homogeneous modification

#### (2) Heterogeneous modification

- Fibres, microfibrils, nanocrystals etc. are modified in a heterogeneous suspension
- → Usually results in surface modification (not necessarily)



# Labelling of hydroxyl groups

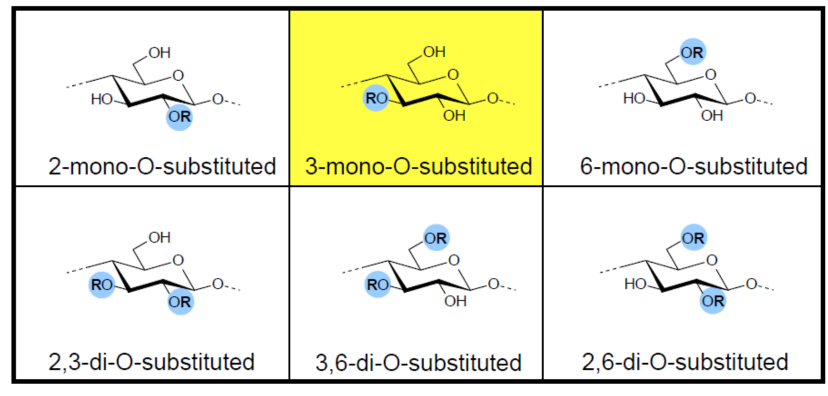


Reactivity in general: 6>2>3



## **Basic concepts**

Regioselectivity: which OH group/groups are selectively modified





## Challenges

- The fundamental challenge in chemical modification of cellulose is that cellulose is relatively inert and does not automatically follow the common rules of organic chemistry
  - Example, cellulose hydroxyl groups are alcohols but they do not form esters with carboxylic acids under normal conditions
- Reproducible and uniform degree of substitution can be difficult to achieve
- Regioselectivity is often difficult to achieve



# Degree of substitution (DS)

- Quality which measures the average amount of substituted hydroxyl groups in an anhydroglucose unit
- Maximum DS is 3: all 3 hydroxyl groups in all anhydroglucose units have been substituted
- On average, if one hydroxyl group per each anhydroglucose unit has been substituted, the DS is 1
- If there is only one hydroxyl group substituted per 10 anhydroglucose units, the DS is 0.1



Cellulose acetate



Carboxymethyl cellulose



Cellulose sulphate



- Cellulose acetate has 39.8% (w/w) acetyl content
- What is the DS?

#### Aids:

Anhydroglucose (inside brackets) M=162 g/mol

Try it out:
We'll come back to this
at the beginning of next
lecture

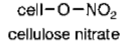
Acetyl group M=43 g/mol

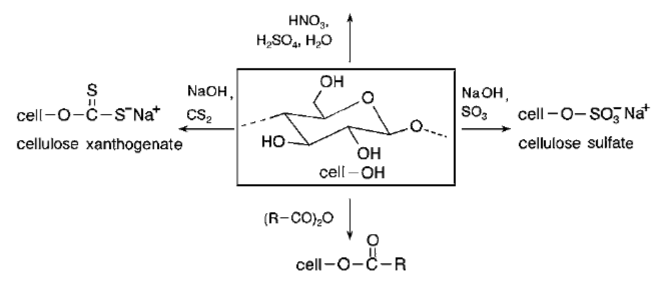


## **Esterification of cellulose**



#### Commercial cellulose esters





R = CH<sub>3</sub>; cellulose acetate (CA)

 $R = CH_3$  und  $CH_2 - CH_3$ : cellulose acetate propionate (CAP)

 $R = CH_3$  und  $CH_2 - CH_2 - CH_3$ : cellulose acetate butyrate (CAB)



# Inorganic cellulose esters

- Cellulose xanthogenate
- Cellulose carbamate
- Cellulose sulphate
- Cellulose nitrate



## Cellulose xanthogenate

Hydroxyl groups are ionized with strong alkali

Half-ester is created by reacting alkoxy cellulose with CS<sub>2</sub>

Cellulose xanthogenate is a half-ester, bearing charge

→ Cellulose xanthogenate is a polyelectrolyte

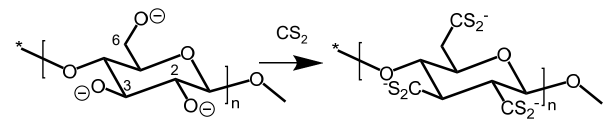


## Cellulose xanthogenate

- The reaction is important in practice because of its use in viscose process
  - Cellulose xanthogenate is produced from native cellulose
  - Cellulose xanthogenate is dissolved
  - The dissolved xanthogenate is regenerated in acid solution, enabling controlled regenaration of cellulose into fibres and films



## Cellulose xanthogenate



Examples of DS and hydroxyl selectivity at different stages of viscose process

Xanthogenated cellulose material	DS at C-2/C-3	DS at C-6
Fiber xanthogenate (DS 0.61)	0.38	0.17
Viscose, non-ripened (DS 0.58)	0.34	0.24
Viscose, moderately ripened (DS 0.49)	0.16	0.32
Viscose, extensively ripened (DS 0.28)	0	0.32



Klemm et al. *Comprehensive Cellulose Chemistry* Vol. 2, Wiley-VCH, 1998.

## Cellulose carbamate

- High temperature reaction (~140°): processed above the melting point of urea
- Catalyzed by metal salts, particularly zinc sulphate is used
- Urea forms isocyanic acid which is the actual reagent with cellulose:

## Cellulose carbamate

OH
$$O + NH_2 - C - NH_2$$

$$R: H \text{ or } -C - NH_2$$

$$R: H \text{ or } -C - NH_2$$

- Cellulose carbamate with DS 0.2-0.3 can be dissolved in aqueous NaOH
- Basis for the CarbaCell process:
  - Aimed at substituting the environmentally hazardous viscose process
  - Enabled by the effortless conversion of carbamate into cellulose in alkali
  - No commercial applications as of yet



## Cellulose sulphate

\* 
$$H_2SO_4$$
 \*  $OHO$   $OH$ 

- Can be prepared in a large variety of systems, usually containing either SO<sub>3</sub> or sulphuric acid
- Water soluble at above DS 0.2-0.3
- Preparation is generally accompanied by severe chain degradation due to acid hydrolysis
- Biomedical applications proposed



#### Cellulose nitrate

- Traditionally produced in a ternary system: HNO<sub>3</sub>/H<sub>2</sub>SO<sub>4</sub>/H<sub>2</sub>O
- Nitrogen contents of commercial cellulose nitrates range from 10.5-13.6%



## Cellulose nitrate

#### **Applications**

- Celluloid (combs, hair ornaments, ping pong balls)
- Explosives (nitrogen content above 12.6%)
- Filters, membranes
- Component in lacquers



# Organic cellulose esters

- Cellulose formate
- Cellulose acetate



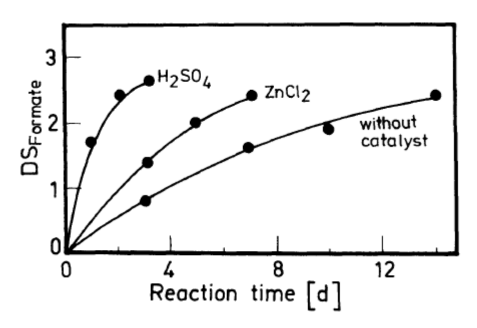
## **Cellulose formate**

- Among the only organic esterifications that proceed spontaneously with the free acid itself
- Cellulose formate is unstable: cellulose formate with DS 2.0-2.5 is decomposed to cellulose and formic acid in 10 h in boiling water



## **Cellulose formate**

#### **Effect of catalyst on formation**



 Very high DS values of cellulose formate can be achieved

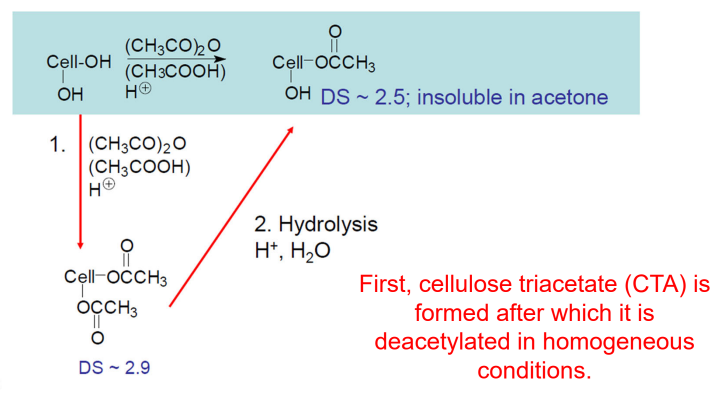


## Cellulose acetate

- Cellulose acetylation proceeds with acetic anhydride and a suitable catalyst in water-free conditions
- Acetic acid alone is not sufficient to acetylate cellulose



# Cellulose acetate (DS 2.5) Commercial synthesis





# Cellulose acetate solubility

#### Strong dependence on DS and position

Liquid	$DS_{Ac}$ range of solubility for partially deacetylated cellulose acetate		
	in C-2/-3/-6 position <sup>a</sup>	in C-2/-3 position <sup>b</sup>	
Water	0.8-1.0	insoluble	
DMF	1.8 - 2.7	1.3-2.8	
Acetone ( $< 0.01 \% H_2O$ )	insoluble	insoluble	
Acetone (1 % H <sub>2</sub> O)	2.3–2.6	2.5–2.6	
Pyridine	0.8 - 2.7	1.2–2.8	
Pyridine/ $H_2O(1:1 \text{ v/v})$	0.6–2.0	1.2–1.6	
Ethyl lactate	1.6–2.7	2.6–2.8	



Klemm et al. *Comprehensive Cellulose Chemistry* Vol. 2, Wiley-VCH, 1998.

#### Cellulose acetate

#### **Applications**

- Coatings (LCD displays)
- Photographic films
- Thermoplastic compounds
- Isolation foils
- Cigarette filters

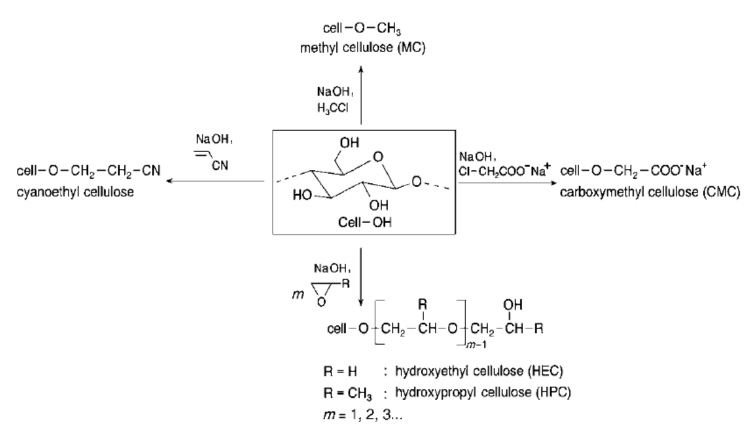


## **Cellulose ethers**

- Methyl cellulose
- Carboxymethyl cellulose



## Commercial cellulose ethers





## Methyl cellulose

Hydroxyl groups are ionized with strong alkali

- Conventional preparation by Williamson reaction with gaseous or liquid chloroform (S<sub>N</sub>2 type nucleophilic substitution)
- 40% NaOH used in the industrial procedure (heterogeneous reaction)
- Methylation performed at ca. 70-90°C

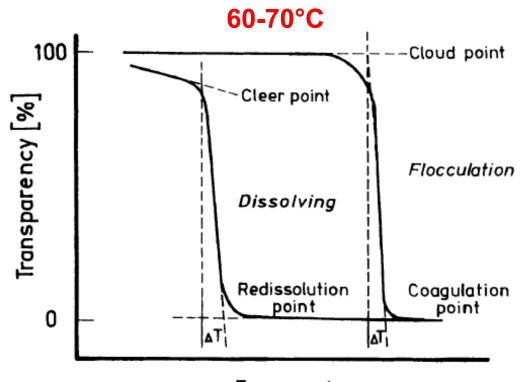


## Methyl cellulose

- DS 1.5-2.0 are produced commercially
- Water soluble at DS 1.4-2.0
- Coordination of water around hydrophobic methyl groups induces water solubility



### Methyl cellulose – thermal response



#### **Between DS 1.7-2.3**

- Solubility is temperature sensitive.
- Gels form above a critical temperature and the gelation is reversible.





Klemm et al. *Comprehensive Cellulose Chemistry* Vol. 2, Wiley-VCH, 1998.

# Methyl cellulose applications

Application area	Proportion (%)
Building industry	47
Dispersion paints	21
Wall paper paints	14
Cosmetics	5
Polymerization	5
Detergents	4
Other	4



 Generally produced by a substitution reaction of monochloroacetic acid to alkoxy cellulose



\* NaOH \* O
$$\ominus$$
 NaOH \* O $\ominus$  O $\ominus$  NaOH \* O $\ominus$  OCH<sub>2</sub>COO- OCH<sub>2</sub>COO- OCH<sub>2</sub>COO-

- 20-30% NaOH concentration
- Temperature 50-70°C
- Exothermic process
- Heterogeneous process in water/isopropanol (or water/tbutanol)



- Commercial grades possess DS values 0.4-0.8
- CMC is water-soluble when DS>0.4
- Aqueous CMC solution does not usually represent a complete dissolution down to the molecular level



- Purified CMC
  - CMC-content min. 98 %

• CMC-content min. 99,5 %

- Technical CMC
  - CMC-content 55 75 %

- Main application areas
  - paper and board
  - oil drilling
  - · paints
  - mining
- Main application areas
  - food
  - dental
  - pharmaceutical
- Main application areas
  - detergent
  - · oil drilling
  - paper and board
  - mining
  - construction



Information from CPKelco

# Regioselective modification of cellulose

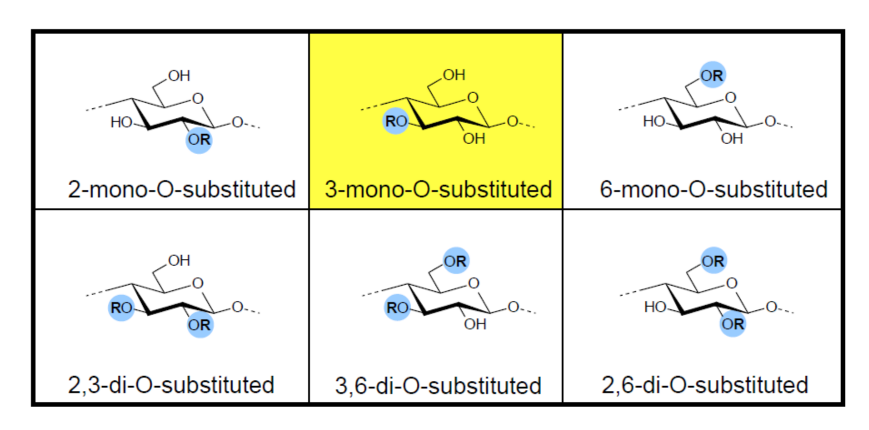


### Regioselectivity issues

- Generally, cellulose hydroxyl groups react in the order O6>O2>O3
- Reactivity of different hydroxyl groups can be tuned by reaction conditions but they are rarely exclusive
- Regioselective synthesis applies various pathways to achieve nearly complete regioselectivity of certain OH group / groups
- Regioselectively prepared cellulose derivatives yield information on the structure-property relationship of polysaccharides and the function of the different hydroxyls on cellulose

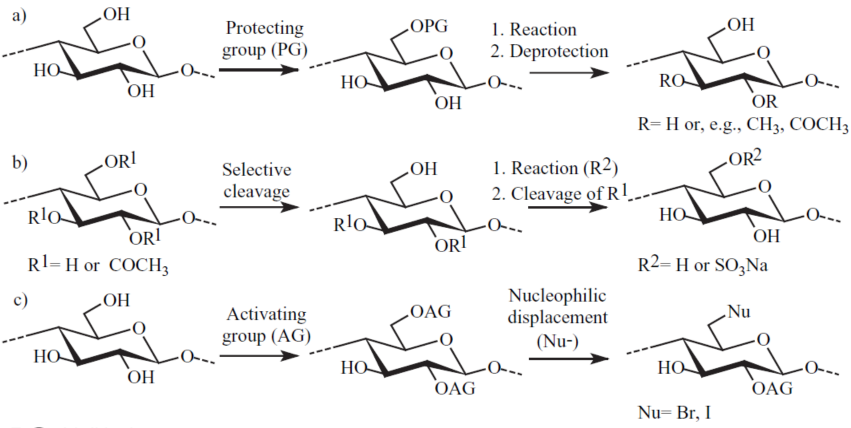


# Regioselectivity nomenclature





# Pathways to regioselectivity





# Protective group: trityl

One of the most popular protective groups for regioselective modification is triphenylmethyl (trityl)



### 6-mono-O-methyl cellulose

OH O 
$$(C_6H_5)_3C_7C_1$$
 OC  $(C_6H_5)_3$   $CH_2=CHCH_2CI$ , OC  $(C_6H_5)_3$  OC



Kondo et al. Carbohydr. Res. 1993, 238, 231.

## **Summary**

- Organic esterification of cellulose requires a more reactive reagent than carboxylic acid, such as acetic anhydride
- Etherification of cellulose proceeds generally via alkoxy ion, generated with harsh alkaline conditions
- Regioselective cellulose modification is a modern trend; it is an important scientific advance in cellulose modification

