Data Article

Thermal, spectroscopic, SEM and rheological datasets of native and quaternized guar gum

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ARTICLE INFO

Article history:
Received 26 March 2020
Revised 4 April 2020
Accepted 6 April 2020
Available online 18 April 2020

Keywords:
TGA/DTA
SEM
NMR
Viscosity

ABSTRACT

The manuscript reports TGA, DTA, SEM and NMR datasets of native and quaternized guar gum as a tool for their characterization. The TGA and DTA data was acquired in temperature range 35–500°C. Based on the TGA experimental values, activation energy plots were drawn to study the stability of native and quaternized polysaccharides (Cai and Bi, 2008 [1]). The data demonstrates the thermal behaviour of quaternized guar gum vis-a-vis native guar gum. The surface morphology of native and quaternized galactomannan was represented by SEM imaging. The 1H–1H COSY was acquired to understand structural changes by quaternization. The rheological measurements of native and modified products were carried out to obtain the viscosity profile of the respective samples. The datasets support the research article ‘Synthesis of quaternized guar gum using Taguchi L (16) orthogonal array’ (Tyagi et al., 2020 [2]).

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https://doi.org/10.1016/j.dib.2020.105565
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**Specifications table**

| Subject | Chemistry |
|---------|-----------|
| Specific subject area | Chemical modification |
| Type of data | Table |
| | Image |
| | Graph |
| | Figure |

**How data were acquired**

- Thermogravimetric: DTG-60 unit (Shimadzu, Japan) under the argon atmosphere.
- NMR analysis: Bruker Avance Neo 500 MHz NMR spectrometer.
- Rheological data: Brookfield DV-III Ultra digital Viscometer.
- SEM imagining: SEM-Zeiss EVO-40 EP. Magnification: up to 10,00,000 X.
- Resolution: 30 nm (HV SE), Carl Zeiss AG Company, Germany.

**Data format**

- Raw and analysed data
  - A. TGA/DTA analysis with activation energy curves of native and quaternized guar gum
  - B. SEM images of native and quaternized gum
  - C. NMR spectra of native and quaternized guar gum
  - D. Viscosity analysis of native gum and quaternized product

**Parameters for data collection**

- TGA/DTA data collection was carried out under the argon atmosphere. The data was collected at a heating rate of 10 °C/min from 35 to 500 °C.
- SEM data was collected by coating the samples with gold
- COSY NMR spectrum was recorded at 125.77 MHz
- Number of scans: 4
- Relaxation delay: 0.5351 s
- Acquisition time: 0.6144 s
- Temperature: 26 °C
- Solvent: D₂O
- Data analysis was processed using iNMR software, version 5.5.5.

**Description of data collection**

- Native guar gum and synthesised derivatives were purified and used for TGA/DTA analysis and NMR spectral recording. The electron microscope imaging and rheological analysis were also performed for these samples.

**Data source location**

- Institution: Forest Research Institute, Dehradun
- Wadia Institute of Himalayan Geology, Dehradun
- Sophisticated Analytical Instrument Facility (SAIF), Punjab University, Chandigarh
- City: Dehradun, Chandigarh
- Country: India

**Data accessibility**

- The raw data files are provided as supplementary files.
- All other data is with this article

**Related research article**

- The data is related to the research article:
  
  - Authors: Rakhil Tyagi, Pradeep Sharma, Raman Nautiyal, Ajeet K. Lakhera and Vineet Kumar. Synthesis of quaternized guar gum using Taguchi L(16) orthogonal array. Carbohydrate Polymers, Volume 237, 1 June 2020, 116136.
  
  - [https://doi.org/10.1016/j.carbpol.2020.116136](https://doi.org/10.1016/j.carbpol.2020.116136)

**Value of the data**

- The datasets show thermal and rheological behaviour of optimized quaternized derivative which will provide understanding for improved utilization of modified products.
- NMR spectral data provided information about structural changes by quaternization of guar gum.
- The surface morphological changes influenced by derivatization is observed in SEM imaging.
- Researchers can use this data to understand quaternization of polysaccharides.
- The data can also be used as reference for future studies on modification of biopolymers.

**1. Data description**

**Images:**
Guar splits were ground to obtain guar gum powder. The powdered guar galactomannan was used as starting material for derivatization by introduction of quaternary ammonium groups as described previously [2].

The thermal stability of native and quaternized guar gum was evaluated by TGA analysis of respective samples. Data was acquired in inert atmosphere (argon) using temperature range 35–500 °C. DTA analysis illustrates the temperature at which maximum weight loss occurred. The analysis was performed using conditions similar to TGA acquisition.

Activation energy plots of the native and quaternized guar gum were derived from TGA data of respective samples by adopting the methodology as described in [1].

Surface morphology of native and modified samples were studied by SEM imaging; data was acquired using magnification range of 1000X–5000X.

COSY spectrum of native and modified samples was recorded by hydrolysis of respective polysaccharide samples followed by removal of acid using rotatory evaporator. Acid free samples were subjected to deuterium exchange and spectral recording was done using D₂O as solvent [2].

Viscosity of 1% (w/v) aqueous solution of native and modified guar gum samples [2] with degree of substitution (DS) ranging from 0.059 to 0.510 was measured by using Brookfield DV-III Ultra Digital Viscometer.

2. Experimental design, materials, and methods

Native and quaternized guar gum samples [2] were dissolved in distilled water and precipitated with methanol using the standard protocol. The precipitated polysaccharide was filtered, washed with aqueous methanol followed by pure methanol and dried in an hot air oven. The pure polysaccharides obtained from both native and quaternized guar gum were used for further analyses.
Fig. 2. Thermo-gravimetric plots: TGA of native (A) and quaternized (B) guar gum; DTA of native (C) and quaternized (D) guar gum.
Fig. 3. Activation energy plots of native (A) and quaternized (B) guar gum.
Fig. 4. SEM images of native (A) and quaternised (B) guar gum at different magnification.
Fig. 5. COSY spectrum of native (A) and quaternized (B) guar gum.
Table 1
Rheological data of native and quaternized guar gum, Spindle 29, Shear rate 2.5 s⁻¹, at 25°C.

| S. No. | Sample code/DS | Viscosity of quaternized guar (Cps) |
|--------|----------------|-------------------------------------|
| 1      | Native guar gum | 6721                                |
| 2      | 1F₁₁F₂₂F₃₃F₄₄F₅₅/0.059 | 2105                                |
| 3      | 1F₁₁F₂₂F₃₃F₄₄F₅₅/0.370 | 5632                                |
| 4      | 2F₁₁F₂₂F₃₃F₄₄F₅₅/0.510 | 2741                                |

2.1. Thermogravimetric analysis

TGA/DTA analysis was carried out using 10 mg of sample in a DTG-60 unit (Shimadzu, Japan) under the argon atmosphere. The scan was carried out at a heating rate of 10°C/min from 35 to 500°C. Data obtained in TGA was further analysed to determine activation energy using Coats and Redfern method [1].

2.2. Scanning electron microscopy

SEM analysis of native and quaternized guar gum sample was recorded using SEM- Zeiss EVO-40 EP instrument with 10,00,000 X magnification and 30 nm resolution. The powdered native and quaternized samples were coated with gold and images were acquired in the magnification range of 1–5 K.

2.3. NMR spectroscopic analysis

The NMR spectra were acquired using a 500 MHz Bruker spectrometer at 25°C. The native and quaternized products were hydrolysed with 0.5 N HCl. Excess of HCl was removed from hydrolysed mass by repetitive co-distillation with methanol using rotary evaporator. The acid-free product was subjected to deuterium exchange (three times). The hydrolysed samples of native and quaternized guar (40 mg) were dissolved in 0.75 mL D₂O (99.95%).

2.4. Determination of rheological properties

The viscosity data of 1% aqueous solution (w/v) of native and quaternized products of DS ranging from 0.059 to 0.510 was determined using Brookfield DV-III Ultra Digital Viscometer. Viscosity of samples was recorded at 25±1°C using spindle 29 at shear rate of 2.5 s⁻¹ (Table 1).

Acknowledgments

The research work reported in the present study has been possible due to financial support provided by Indian Council of Agricultural Research (ICAR), New Delhi to work on the project entitled ‘Value addition to Guar Gum and its by products’ under Consortium Research Project on Secondary Agriculture being implemented by Central Institute of Post Harvest Engineering and Technology (CIPHET), Ludhiana as the Lead Institute. The authors are thankful to the Director, Forest Research Institute (FRI), Dehradun for making available the wet laboratory and other vital research facilities. The authors sincerely acknowledge Wadia Institute of Himalayan Geology, Dehradun for recording SEM data and Sophisticated Analytical Instrumental Facility (SAIF), Punjab University, Chandigarh for spectral data recording.
Conflict of Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.dib.2020.105565.

References

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