Thermoanalytical studies (TG–DTG–DSC, Py–GC/MS) of sodium carboxymethyl starch with different degrees of substitution

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Abstract
The paper presents the results of thermal analysis of polymer material in the form of starch derivatives in the form of sodium carboxymethyl starch (CMS–Na) with degree of substitution (DS) in the range 0.2–0.9 for the preparation of foundry binder. In this work, the thermal behavior of the modified starch and qualitative assessment of degradation products released during pyrolysis were determined and comprised. The analysis of the course of progressive decomposition of the starch material under controlled heating in the range of 25–1000 °C in anaerobic atmosphere was based on the results of thermal analysis methods (TG–DTG–DSC) in combination with the results of pyrolysis–gas chromatography–mass spectrometry (Py–GC/MS). The detailed TG–DTG–DSC analysis allowed to determine and compare the temperature at which the process of decomposition of carboxymethyl starches sodium salts with different degrees of substitution begins and to determine the course of its degradation under conditions corresponding to the contact of the foundry binder in the form of starch material with liquid metal (conditions like in foundry mold). Thermogravimetric analysis shows that decomposition processes are multistage, and dehydration is the first step of decomposition. Moreover, TG–DTG–DSC analyses indicate that the thermal stability and the decomposition path of tested compounds depend on the DS. Results of Py–GC/MS studies showed that the formation of decomposition products (including cyclic and aromatic hydrocarbons) in a predetermined temperature range is lower in the case of CMS–Na with high DS.

Keywords Polymer · Sodium carboxymethyl starch · TG–DTG–DSC · Py–GC/MS · Thermal degradation · Foundry binders

Introduction
In the foundry industry in the production cycle, it is possible to use organic, inorganic and mineral substances, which, as binding materials, ensure obtaining durable casting forms based on the mineral matrix, at the same time giving the possibility of making castings with the reduction in their defects. Binders in sand molds are a key component determining their properties and produced castings quality by the binding strength of the of the matrix grains of sand. However, some of these materials may be harmful to the environment during the technological process, even in their initial form. Therefore, a lot of research work is focused on the modification of materials or the development of new substances toward the development of environmentally friendly casting technologies [1–5]. The scientific work on the development of non-toxic and degradable binders also seems to be stimulated by the progress in research on modification of natural polymers, e.g., native starch, which has been observed in recent years. Poor properties of bonding sand by native starch-based binders show that they are not a suitable alternative to the currently used commercial binding materials from technological reasons [6, 7]. But many works carried out in the field of application of natural polymers in foundry materials engineering prove that it is the modified form of starch that can provide a certain alternative to harmful components currently used in mold technology, while at the same time ensuring that molding sands with appropriate strength and

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performance parameters are obtained [2, 3, 5, 8]. Despite the constantly increasing possibilities of using starch derivatives, many aspects related to the process of their obtaining, crosslinking and degradation have remained unexplained. Moreover, the influence of the modification method, e.g., on the thermal properties of binders in the form of modified starch in relation to the application in foundry processes, is still studied.

In our work, the sodium carboxymethyl starch (CMS–Na), obtained by potato starch etherification, was chosen as the subject of research. During the chemical modification of natural starch, hydroxyl groups with carboxymethyl groups (with hydrophilic properties) are substituted in the rest of the glucopyranose rings. The number of substituted hydroxyl groups can be controlled quantitatively, and the effect of the modification process is characterized by the degree of substitution (DS) [9, 10]. It results from the literature research that the hydrophilic group (with –COONa) share in the CMS–Na stems results in beneficial changes in the properties of the modified starch obtained. For example, the process of starch carboxymethylation already ensures its solubility in cold water and at the same time increases adhesion properties [11–14].

Due to the degree of substitution, the sodium carboxymethyl starch is classified into three groups, as starch derivatives with low degree of substitution: DS < 0.3; with medium degree of substitution: DS in the range of 0.3–0.7; with high substitution degree: DS in the range of 0.7–1.4 [15–18].

Results of research on foundry applications of sodium carboxymethyl demonstrated that with increasing DS of modified starch, it is possible to increase the strength of molds and its other properties [2, 6, 19, 20]. In our own research, CMS–Na was successfully used as a water-soluble binder for molding sands susceptible to both chemical and physical crosslinking (by conventional heating and exposure to electromagnetic radiation) [5, 6, 8].

From the point of view of pouring a liquid alloy casting into foundry sand molds, important issue is it is to determine the thermal stability of a binder during process. Namely, course of thermal decomposition of the organic binder in molding sand determines the result in the formation of significant casting defects, knock-out properties or adversely affect the efficiency of thermal reclamation methods of used molding sands [21–25]. In terms of environmental protection and the assessment of the binder’s harmfulness, it is also important to determine the quality (composition) of the decomposition products of the binding material released during rapid heating. In this paper, the subject matter related to the explanation of relations between substitution degree and CMS–Na properties in the context of recognition of the degradation process in the binder-sand system subjected to interaction with liquid metal has been discussed. In this work, efforts were made to supplement the knowledge by thermal analysis (TG–DTG–DSC) and pyrolysis–gas chromatography–mass spectrometry (Py–GC/MS) CMS–Na with the degree of substitution in the range of 0.20–0.87 as a main binder in the molding sand technology.

**Experimental**

**Materials**

The following samples of CMS–Na with various DS were subjected to thermoanalytical studies:

- Low-substituted CMS–Na_L with DS 0.2; modification by semi-dry method [2],
- Medium-substituted CMS–Na_M with DS 0.5; modification by semi-dry method [2, 12, 26],
- High-substituted CMS–Na_H with DS 0.9; modification by slurry method—one step synthesis [11, 12].

Moisture content of the starting product was 7.0–8.0%.

**Methodology**

**Methods of thermal analysis (TG–DTG–DSC)**

Thermal analysis was performed by using a thermal analyzer NETZSCH STA 449 F3 Jupiter® design to simultaneous thermogravimetric (TG), differential thermal analysis (DTG) and differential scanning calorimetry (DSC) measurements, which ensure to obtain independent signals recorded under the same measurement conditions: heating rate, atmosphere and pressure. This method of analysis provided a higher efficiency and allowed obtaining comprehensive information on the thermal characteristics of the samples of tested starch derivatives. The measurement was performed under a nitrogen gas atmosphere in a temperature range of 25–1000 °C at a gas flow rate of 40 mL min⁻¹. The rate of heating of the test sample was 10 °C min⁻¹.

**Pyrolysis–gas chromatography–mass spectrometry (Py–GC/MS)**

The CMS–Na samples (2–4 mg) were thermally degraded in Pyroprobe 5000 pyrolyser (CDS Analytical) in an inert atmosphere (helium). The sample was heated to a set temperature at a rate of 10,000 °C/s for a time interval of 3–4 s. Then chromatographic separation (Focus GC, Thermo Scientific) was carried out according to the assumed measurement conditions: Initial heating at 40 °C lasted for 3 min, then the temperature was increased to 100 °C (at a rate of 3 °C min⁻¹) and heated for 3 min. Products of pyrolysis were separated on a non-polar TR-SQC tested column (30 m
length, 0.25 mm diameter). The next temperature increase to 250 °C was carried out at the rate of 20 °C min⁻¹, and the sample was heated for 3 min at the rate of helium flow in the column of 1 mL min⁻¹ and at the division of the sample at the ratio of 1:30. The injector was maintained in the SPLIT mode. The last stage of the study included detection (ISQ, Thermo Scientific) according to the following parameters: ion source temperature 250 °C, electron energy 70 eV at full range m/z. The qualitative analysis was performed with the use of Xcalibur 2.2 software based on the NIST MS Search 2.0 database as well as on mass spectral data and retention time comparisons reported in literature.

Results and discussion

Thermal analysis TG–DTG–DSC

Series of TG, DTG and DSC curves obtained in anaerobic atmosphere for CMS–NaL, CMS–NaM and CMS–NaH samples are shown in Figs. 1–3.

The parameters that characterize the thermal decomposition are as follows: mass loss (Δm), the temperature of maximum rate of mass loss (T_{peak}), temperature corresponding to 10% mass loss (T_{10}) and the temperature corresponding to 50% mass loss (T_{50}), temperature of exo- or endothermic effects occurring in selected temperature range (these parameters are presented in Table 1). On the course of thermal curves, it is possible to determine the subsequent stages of degradation of the binder.

![Fig. 1 TG curves of CMS–Na with different degrees of substitution](image1)

![Fig. 2 DTG curves of CMS–Na with different degrees of substitution](image2)

![Fig. 3 DSC curves of CMS–Na with different degrees of substitution](image3)

The changes in the material due to the rising temperature for CMS–Na samples are multistage (Figs. 1–3). Three mass losses were recorded on the TG curves of CMS–NaL: 10.7%, 55.7% and 21.1% at temperature 63 °C, 280 °C and 871 °C at which the mass loss rate is maximum (based on DTG curve) and CMS–NaM: 12.2%, 51.7% and 26.1% with T_{peak} at 64 °C, 291 °C and 890 °C, respectively. Thermal decomposition of CMS–NaH was more complex as it was decomposed with four mass losses of 15.1%, 3.6%, 17.0% and 15.2% with mass loss rate maximum at 45 °C, 291 °C, 733 °C, 861 °C, respectively. The degradation process of the
sodium salt of carboxymethyl starch begins at temperature above ~ 220 °C. All tested samples in an anaerobic atmosphere were not completely decomposed, and their residue determined at 1000 °C was 12.5%, 10.0% and 9.1% for CMS–NaL, CMS–NaM and CMS–NaH, respectively.

To evaluate and compare the thermal stability of CMS–Na, the parameters $T_{10}$ and $T_{50}$, respectively, were considered. In the case of CMS–NaL, mass loss of 10% is recorded at temperature of 175 °C; comparative measurements to CMS–NaM ($T_{10} = 132$ °C) and CMS–NaH ($T_{10} = 102$ °C), were made. On the other hand, mass losses of 50% take place until a temperature of 306 °C is attained in the case of CMS–NaL, 309 °C for CMS–NaM, and 372 °C for CMS–NaH, respectively. Analyzing these mass losses by thermal decomposition, it can be affirmed that the CMS–NaH presents a highest thermal stability (higher than that of CMS–NaL and CMS–NaM). The above-mentioned parameter $T_{50}$ confirmed that CMS–NaH has higher thermal stability than in the case of CMS–Na with low and medium substitution degrees; however, this statement should be applied to the temperature range of ~700–720 °C, because observing further changes in the sample above, it was found that CMS–NaH is less stable in the range of 730–1000 °C.

On the DSC curves, as shown in Fig. 3 two types of the thermal signals were clearly visible. Three endothermic effects at 63 °C, 752 °C and 881 °C and three exothermic effects at 291 °C, 792 °C, 881 °C were recorded for the CMS–NaL sample. CMS–NaH mass losses were accompanied by two endothermic effects at 723 °C and 861 °C and two exothermic effects at 792 °C, 861 °C visible as a DSC curve signal.

It was found that the first loss of mass of three CMS–Na samples in the temperature range 25–220 °C was caused by evaporation of water absorbed on the granules of the starch product and dehydration of the material above 100 °C (the small endothermic peaks on DSC curves indicate this).

The second significant loss in the temperature range ~220–550 °C was attributed to the fragmentation of polymer

| Sample   | Temperature range/°C | Mass loss/Δm/% | $T_{	ext{peak}}$/°C | Thermal effect peak/°C | $T_{10}$/°C | $T_{50}$/°C |
|----------|----------------------|----------------|---------------------|------------------------|-------------|-------------|
| CMS–NaL  | 25–215               | 10.7           | 63                  | 63/endo                | 175         | 306         |
|          | 215–536              | 55.7           | 280                 | 291/exo                |             |             |
|          | 536–1000             | 21.1           | 871                 | 752/endo               | 792/exo     | 881/endo    |
|          |                      |                |                     |                        |             |             |
|          | Residue: 12.5% (at 1000 °C) |              |                     |                        |             |             |
| CMS–NaM  | 25–216               | 12.2           | 64                  | 64/endo                | 132         | 309         |
|          | 216–527              | 51.7           | 291                 | 291/exo                |             |             |
|          | 527–1000             | 26.1           | 890                 | 733/endo               | 762/exo     | 881/endo    |
|          |                      |                |                     |                        |             |             |
|          | Residue: 10.0% (at 1000 °C) |              |                     |                        |             |             |
| CMS–NaH  | 25–221               | 15.1           | 45                  | –                      | 102         | 372         |
|          | 221–556              | 43.6           | 291                 | 291/exo                |             |             |
|          | 556–763              | 17.0           | 733                 | 723/endo               |             |             |
|          | 763–1000             | 15.2           | 861                 | 792/exo                | 861/endo    |             |
|          |                      |                |                     |                        |             |             |
|          | Residue: 9.1% (at 1000 °C) |              |                     |                        |             |             |

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**Table 1** Thermoanalytical data (TG, DTG, and DSC) for CMS–Na samples

- **a** Based on TG curves
- **b** Based on DTG curves
- **c** Based on DSC curves
Thermoanalytical studies (TG–DTG–DSC, Py–GC/MS) of sodium carboxymethyl starch with different…

chains, cracking of C–O–C bonds in the glycosidic bond (bonds between starch repeat unit in polymer chain), opening of glucopyranose rings and formation of volatile compounds (e.g., H₂O, CO, CO₂), as confirmed by literature data [27–29].

Irreversible changes in the structure of the CMS–Na sample were recorded at 280–290 °C (in this temperature range the maximum rate of changes and a strong exothermic effect were observed; moved to higher temperature range if DS was higher). The literature data show that the thermal effect on DSC curves recorded for CMS–Na samples at ~ 300 °C is characteristic for natural and modified polysaccharides and is attributed to the formation of volatile products in the process of thermal degradation. Thermal reactions for starch start around 300 °C with thermal condensation between hydroxyl groups of starch chains to form ether segments and liberation of water molecules and other small molecular species. Dehydration of neighboring hydroxyl groups in the glucose ring also occurred, resulting in the formation of C=C bonds or breakdown of the glucose ring [28, 30–32].

The next large loss of mass up to ~530–550 °C for CMS–Na samples was connected with detachment of side groups and further cracking of weak glycosidic bonds. In this temperature range of 220–1000 °C, the mass losses may be associated also with much efficient course of material degradation and detachment of the side groups –COONa (from substituted groups). The endothermic peaks visible on DSC curves at higher temperature were connected to the breaking of the bonds in the structure of samples and may be associated with a carboxy combustion in an anaerobic atmosphere.

Based on obtained results, it was found that the DS of CMS–Na—average amount of substituted OH groups in polymer chains—may influence the course of starch decomposition in tested samples, which was also confirmed in other research works in natural and modified polymer systems [30, 31, 33, 34]. So, modified structure and the presence of sodium cation in CMS–Na can play a key role in the thermal processes of high-substituted modified starches, increasing the degradation within a certain temperature range (above 730 °C), as has also been observed in other natural materials [30, 33, 35]. In addition, at 1000 °C, the residual mass of CMS–Na samples probably contained carbonized carbon and Na₂CO₃ (which formation begins in an anaerobic atmosphere in the temperature range of approx 600–750 °C [36]).

Table 1 summarizes the characteristic thermal data selected from the TG–DTG–DSC curves for CMS–Na samples.

### Analysis of products decomposition

Based on data obtained from TG–DTG–DSC tests, qualitative analysis of decomposition products released at a selected temperature was performed for CMS–Na samples using the method of pyrolytic gas chromatography coupled with mass spectroscopy (Py–GC/MS). Measurements were conducted at 300 °C (temperature of strongest exothermic effect), 550 °C (approximate temperature between the second and third mass loss, see Fig. 1) and 1000 °C (the maximum temperature). Figures 4–6 show the Py–GC/MS results of the CMS–NaL, CMS–NaM, CMS–NaH samples (for retention data and peak identification, see Table 2).

On the basis of the obtained results, it was found that at 300 °C the decomposition of CMS–Na samples took place with the separation of a few compounds registered with the chosen measurement method (Table 2). It was noted that the

![Py–GC/MS chromatograms of CMS–NaL under the different pyrolysis temperatures](image)
signal from these compounds was more intense for samples with lower DS and there were more of them. Small peaks on CMS–Na_L and CMS–Na_H chromatograms were identified as CO_2. The course of the CMS–Na_H chromatogram indicates that the share of CO_2 in degradation products is unnoticeable. Water could be the main product of decomposition below about 300 °C, and H_2O is formed by inter- and intramolecular condensation of hydroxyl groups in starch chains [11, 37, 38]. So, taking into account the results of thermal analysis indicating a significant mass loss in the range above ~220 °C, with maximum rate at 300 °C, it was found that pyrolysis in 300 °C could occur with the release of H_2O. However, the column type used in the study does not allow detecting signals from the polar components of the sample decomposition.

The release of cyclic hydrocarbons in the form of, e.g., cyclopenta-1,3-diene, 2-methylcyclopentane-1-one, cyclopent-2-en-1-one and 2,4-dimethylcyclopentane-1,3-dione was observed on chromatograms of CMS–Na_L sample pyrolyzed at 550 °C. Aromatic hydrocarbons were also formed by decomposition of the low-substituted CMS–Na at 550 °C in the form of benzene and its derivatives (Table 2). No cyclic or aromatic hydrocarbons were detected in the decomposition products CMS–Na_M and CMS–Na_H at this temperature.

The identification of carbon dioxide at 1000 °C indicated the progressive process of decomposition. The presence of benzene, toluene, styrene, phenol and/or their derivatives was found in all CMS–Na degradation products at 1000 °C. Significant intensification of CMS–Na_H sample decomposition with formation of more degradation products took place at temperatures above 550 °C. At 1000 °C, less number of groups of aromatic hydrocarbons was formed in comparison with CMS–Na_L and CMS–Na_M, which was caused by higher amount of sodium on the course of formation of this type of compounds.

The Py–GC/MS results imply that the increased DS of the samples (polymer with increased content carboxymethyl group with Na⁺) alters the decomposition course. This conclusion is confirmed by the decreasing amount of the decomposition products with increasing degree of substitution of CMS–Na.
Conclusions

In this paper, thermal degradation properties and effect of pyrolysis of carboxymethyl starch with different DS were analyzed and compared:

1. The results of the TG–DTG–DSC analyses of starch binder CMS–Na show that its thermal decomposition was a multistage process. Degradation of the binder begins at ~220 °C. The subsequent mass losses at temperature range 25–1000 °C are determined as evaporation of water adsorbed in molecule, sodium carboxymethyl starch dehydration, next breakage of glycosidic bonds in the polymer chains and intensify the fragmentation of chain.

2. Varied thermal behavior was observed for starch samples at temperature above to 220 °C due to their difference in structural modification. CMS–Na with high DS is more

Table 2 Results of Py–GC/MS measurements for CMS–Na samples at different temperatures

| Peak no. | Compound (IUPAC name) | No. CAS | Molecular formula | Molecular weight/u | Retention time \(R_t/\text{min} \) |
|----------|-----------------------|---------|-------------------|--------------------|-----------------------------|
|          |                       |         |                   | CMS–Na\(_L\)       | CMS–Na\(_M\)               | CMS–Na\(_H\)               |
| 1.       | Carbon dioxide        | 124-38-9| CO\(_2\)          | 44                 | 2.16\(^a\)                 | 2.18–2.46\(^a\)           | 2.16\(^a\)               |
|          |                       |         |                   | 2.17\(^b\)         | 2.16\(^b\)                 | 2.15\(^b\)               |
|          |                       |         |                   | 2.16\(^c\)         | 2.16\(^c\)                 | 2.13\(^c\)               |
| 2.       | Cyclopenta-1,3-diene  | 542-92-7| C\(_3\)H\(_6\)    | 66                 | 2.71\(^b\)                 | 2.72\(^c\)               | 2.72\(^c\)               |
|          |                       |         |                   | 2.68\(^c\)         | 2.68\(^c\)                 | 2.68\(^c\)               |
| 3.       | Butane-2,3-dione      | 431-03-8| C\(_2\)H\(_4\)O\(_2\) | 86                 | 3.64\(^c\)                 | 3.51\(^c\)               | 3.49\(^c\)               |
| 4.       | Benzene               | 71-43-2 | C\(_6\)H\(_6\)    | 78                 | 4.57\(^b\)                 | 4.46\(^c\)               | 4.44\(^c\)               |
|          |                       |         |                   | 4.46\(^c\)         | 4.46\(^c\)                 | 4.46\(^c\)               |
| 5.       | (1E)-buta-1,3-dien-1-ol| 70411-98-2| C\(_4\)H\(_6\)O | 70                 | 5.02\(^c\)                 | –                         | –                         |
| 6.       | Pentane-2,3-dione     | 600-14-6| C\(_4\)H\(_6\)O\(_2\) | 100                | 5.60\(^b\)                 | 5.53\(^c\)               | 5.52\(^c\)               |
| 7.       | Acetic acid           | 64-19-7 | C\(_2\)H\(_4\)O\(_2\) | 60                 | 5.86\(^b\)                 | 5.82\(^c\)               | –                         |
|          |                       |         |                   | 5.86\(^b\)         | 5.82\(^c\)                 | 5.82\(^c\)               |
| 8.       | 1-Hydroxypropan-2-one | 116-09-6| C\(_3\)H\(_6\)O\(_2\) | 74                 | 6.43\(^b\)                 | 5.79\(^b\)               | –                         |
|          |                       |         |                   | 6.43\(^b\)         | 6.19\(^c\)                 | 6.19\(^c\)               |
| 9.       | Toluene               | 108-88-3| C\(_6\)H\(_6\)    | 92                 | 7.36\(^c\)                 | 7.3\(^c\)                | 7.29\(^c\)               |
| 10.      | 1-Hydroxybutan-2-one  | 5077-67-8| C\(_4\)H\(_8\)O\(_2\) | 84                 | 9.73\(^b\)                 | 9.49\(^b\)               | –                         |
|          |                       |         |                   | 10.41\(^c\)        | 9.59\(^c\)                 | 9.59\(^c\)               |
| 11.      | Cyclopentanone        | 120-92-3| C\(_5\)H\(_8\)O    | 106                | 11.4\(^c\)                 | 11.36\(^c\)              | –                         |
| 12.      | Ethylbenzene          | 100-41-4| C\(_8\)H\(_10\)   | 106                | 11.57\(^c\)                | 11.53\(^c\)              | 11.52\(^c\)              |
| 13.      | 1,2-Xylene            | 95-47-6 | C\(_8\)H\(_10\)   | 106                | 12.47\(^b\)                | –                         | 12.27\(^c\)              |
| 14.      | 2-Methylcyclopentan-1-one | 1120-72-5| C\(_8\)H\(_10\)O\(_2\) | 98                | 12.36\(^b\)                | 12.36\(^b\)              | –                         |
| 15.      | Ethynylbenzene        | 536-74-3| C\(_6\)H\(_6\)    | 102                | 13.41\(^b\)                | 13.34\(^b\)              | –                         |
| 16.      | Styrene               | 100-42-5| C\(_8\)H\(_8\)    | 104                | 13.61\(^c\)                | 13.56\(^c\)              | 13.54\(^c\)              |
| 17.      | Cyclopent-2-en-1-one  | 930-30-3| C\(_5\)H\(_8\)O    | 82                 | 14.19\(^b\)                | 13.97\(^c\)              | –                         |
| 18.      | 1-Ethyl-2-methylbenzene| 611-14-3| C\(_9\)H\(_12\)   | 120                | 14.03\(^c\)                | 14.03\(^c\)              | –                         |
| 19.      | 2-Methylcyclopent-2-en-1-one | 1120-73-6| C\(_8\)H\(_10\)O\(_2\) | 96                | 16.85\(^c\)                | 16.81\(^c\)              | 16.8\(^c\)               |
| 20.      | 1-Ethenyl-2-methylbenzene | 611-15-4| C\(_8\)H\(_10\)   | 118                | 18.44\(^b\)                | 18.63\(^c\)              | –                         |
|          |                       |         |                   | 18.68\(^c\)        | 18.68\(^c\)                | 18.68\(^c\)              |
| 21.      | Phenol                | 108-95-2| C\(_6\)H\(_4\)O    | 94                 | –                           | 19.9\(^c\)               | –                         |
| 22.      | 1-Ethynyl-3-methylbenzene | 766-82-5| C\(_8\)H\(_8\)    | 116                | 22.50\(^c\)                | 22.52\(^c\)              | 22.49\(^c\)              |
| 23.      | 2,4-Dimethylcyclopentan-1,3-dione | 34598-80-6| C\(_4\)H\(_10\)O\(_2\) | 126                | 23.77\(^b\)                | –                         | 23.75\(^c\)              |
| 24.      | 3-(2-Methyl-2-propanyl)-2,5-furandion | 18261-07-9| C\(_8\)H\(_10\)O\(_3\) | 154                | 24.14\(^c\)                | 24.09\(^c\)              | –                         |

\(^a\)Compound detected in 300 °C
\(^b\)Compound detected in 550 °C
\(^c\)Compound detected in 1000 °C
stable in temperature up to ~730 °C than CMS–Na samples with low and medium DS; however, above this temperature the degradation is more intense. At 1000 °C, the samples are not completely decomposed (the remaining mass may stand carbonized carbon and Na2CO3).

3. The thermal decomposition of CMS–NaA, CMS–NaB, CMS–NaM, CMS–NaF samples was studied by Py–GC/MS analysis which showed that the chemistry of their degradation was different depending on the degree of substitution. It was shown that the amount of substituted groups in CMS–Na polymer chains (with sodium ion presence) has a significant impact on course of CMS–Na degradation and decreases the tendency to formation of CO2, cyclic and/or aromatic hydrocarbons in the range of 300–1000 °C for samples with high DS. Therefore, it has been proved that starch etherification does not adversely affect the ecological context of the molding sands under consideration with the participation of CMS–Na.

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