Measurement of Molding Sand Elasticity

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Abstract
The progressive mechanization and automation of industrial equipment is the driving force of progress, not only in the field of production but also in the measuring and control equipment. In mold production, the automation of processes such as forming molds and cores along with their assembly has led to increases in serial production, reductions in defects, and the shortening of molding times, among others. Thanks to automation in mold and core departments and the use of all sorts of manipulators, mold production in foundries has gained momentum.
Unfortunately, in addition to the mentioned advantages, there are also new challenges as to the quality and properties of the molding and core sands used in highly automated foundries. This article presents recent research on molding sand elasticity. The topic was introduced as an attempt to answer the new needs of highly mechanized foundries.
The article discusses a new method of measuring the resistance of molding materials to undergoing mechanical deformation (molding sand elasticity), with an additional analysis of the bending strengths of the tested samples. Precise measurements, test sample preparation, and interpretation of the received results are presented in the article.

Keywords: molding sands, mechanical deformation, elasticity, hot-distortion

1. NEW CHALLENGES

Cores and molds are subjected to a variety of mechanical and thermal factors at each stage of their production and use. There are various technological operations that involve many possible sources of damage to the molds and cores. Operations such as the removal of cores from core boxes, removal of patterns from molds, core and mold assembly, applying weights to molds, etc. can generate cracks in cores and molds that may later lead to defects in the castings (Figs. 1 and 2). This is true especially in highly mechanized foundries, where such operations are often performed by manipulators. The reason for this is the high brittleness of the cores and molds. Hence, modern technology is looking for molding mixtures with relatively high flexibility/elasticity. This mainly applies to core sands prepared with the addition of synthetic resins [1–4].

When cured, the synthetic resins used in the preparation of molding and core sands can be classified into polymer groups. The polymeric materials sustain different proportions of both elastic deformation and elastic and plastic (permanent) deformation during handling. The shares of the deformations depend on many parameters; for example, strain rate, temperature, deformation range, and the type of bond formed in the polymer material [5–8].

Both foundry molds and cores are subjected to many destructive factors during their execution and assembly. In order to not damage the cores and molds, the used molding sand must not be too brittle to be damaged; however, at the same time, it must retain its shape and stiffness.
The presented device performs measurements in two modes:

- DMA – thermal deformation (hot distortion parameter),
- DE – measurement of elasticity (registration of deflection arrows with force registers).

Thanks to its modern and compact design as well as its use of new measuring systems, the device provides accurate results. The included software enables the export of results to popular office systems (Excel) and archives them in an independent database regardless of the device status.

2.1. Molding sand mechanical deformation – elasticity

In the study of polymer materials [7–12], elasticity is defined as a property that allows to make reversible shape changes under the influence of external forces. It should also be noted that, as a rule, we are not dealing with materials showing only one kind of deformation. In foundry molds, both spring and plastic deformation can occur in various shares. They may depend on different parameters; e.g., temperature or deformation speed.

The DE-module, dedicated to mechanical deformation measurement, is used for estimating molding sand elasticity. It gathers information about the development of force over time with the simultaneous registration of the bend extend. This information is necessary for estimating the maximum bending force and maximum bending radius.

The elasticity measurement is based on the analysis of the deflection pattern of a standard longitudinal fitting during bending. The tooling (Fig. 4) allows us to perform measurements of the indentation force over time while recording its displacement. This allows the user to determine the bending strength ($R_g$) and deflection arrows [13]. The details concerning the elasticity measurement are described in earlier works [14].

The apparatus allows us to perform measurements of the indentation force over time while recording the indenter displacement. This allows the user to determine the bending strength of the sample as well as the deflection arrows.
The software provided by the manufacturer allows us to view the results in real time. In addition, it has the ability to display ready-made graphs showing the relationship between the applied force versus time as well as displacement and displacement as a function of time (Fig. 5).

By analyzing the graphs obtained by means of the device, it can be seen that the point of maximum force is applied at the moment of fracture under the indentation, while the part of the graph after that point is crossed is connected with the indenter retrieving back to the starting point of the measurement.

The program records the most important data such as time of measurement, indenter movement speed, maximum and momentary displacement and indenter force, sample size, and name.

The force-measuring range is 0 to 900 N; in addition, it is possible to adjust the indenter’s movement speed from 0 to 70 mm/min in increments of 1 mm. The built-in database allows us to determine the size of the fitting for each type of strength test (from \( R^1_g \) to \( R^9_g \)), and it is possible to enter any dimension of the test piece.

There are two types of supports (20 mm and 38 mm), which can be mounted in two spacings – 10 and 15 cm.

Thanks to this, the measurement capabilities of the device increase significantly.

The bending strength tests were carried out on standard longitudinal samples after 24 h of curing [13].

### 2.2. Molding sand thermal deformation – hot distortion parameter

The hot distortion parameter tests were carried out on numerous devices [13, 15–20]. In this research, a new approach to this measurement was used. The measurements of the thermal deformation of the molding sands were carried out on rectangular samples measuring 114 × 25.4 × 6.3 mm [13].

One of the ends of the fitting is fixed in the jaws of the device (Fig. 6), while the tilt sensor rests on the opposite (free) end of the sample. In addition, a temperature sensor is provided parallel to the fitting to increase the accuracy of the temperature measurement [13].

The heating system of the device allows the heating temperature to be adjusted independently for the upper and lower heating elements (which are \( 2 \times 400 \) W ceramic heaters, each with an independent built-in thermocouple).

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**Fig. 5.** Elasticity curves: a) deformation \( L \) (brown curve) and applied force \( F \) (orange curve) as a function of time; b) applied force \( F \) as a function of deformation \( L \) (red curve), deflection arrow (blue curve)

**Fig. 6.** LRu-DMA measuring device from MULTISERW-Morek Company – hot distortion module

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The device also allows the heating power to be controlled from 0 to 100% of maximum power (with increments of 1%). The heating temperature ranges from room temperature to 900°C and can be set at 1°C increments, while the maximum deformation can range up to 10 mm. In addition, metal overlays can be used to limit the heat emission field.

There is also the possibility of heating the upper and/or lower heating element to a set temperature (power %) and placing it over and/or under the fitting. This gives us the opportunity to analyze changes in molding sand deformation, both in the case of the sudden exposure of the molding mixture to high temperatures (contact with the liquid alloy) and slow heating (heat radiation from the surface of the liquid metal) [21].

3. OWN RESEARCH

The following article focuses on the deformation of molding samples under pressure of force. The influence of the type of binding agent on the elasticity as well as the morphology of the crack surface are also taken into account. Considering the differences that occur between the different molding technologies, the type of bond destruction can play a leading role in molding sand deformation (among other properties) [22–25].

The presented results are a continuation of the authors’ ongoing research described in detail in previous works [26–30].

The research was carried out in the laboratory of the Department of Molding Materials, Mold Technology, and Cast Non-Ferrous Metals at the Faculty of Foundry Engineering at AGH.

Both organic and inorganic binders were chosen for the study. They can be characterized by different mechanisms of destruction [27, 28]. Organic bonded molding sands can be characterized by adhesive destruction, while inorganic molding sands feature cohesive destruction [31]. The mechanisms of the destruction of these molding sands were studied in detail in previous works [32, 33].

As author Stanisław Dobosz [32, 33] points out in his thesis, the destruction of inorganic molding sands runs through the binder. This means that the adhesion forces between the binder and the grain surface are greater than the cohesion forces, which promotes breakage at the weakest place inside the binder layer (Fig. 7).

All of the molding sands that were chosen for this research were prepared according to the manufacturer instructions using standard compositions (Tab. 1). Four representative molding mixtures that are widely used in the foundry industry were chosen for the research [21].

The following resins/binders were used:

- sands manufactured in cold-box technology – GASHARZ 6966 and AKTI-VATOR 7624 [34],
- self-hardening molding sands with furfuryl resin – XA-20 resin and 100T hardener [34],
- alkdy resin technology – SL2002 resin and KL1 catalyzer [35],
- inorganic molding sands – 145 hydrated sodium silicate and Flodur3 [14, 21, 35].

Quartz sand was used in all of the conducted tests. According to Polish standard PN-85/H-11001, it classifies the tested sand as medium. In the studied matrix, the value of the main fraction is 84%, which determines the sand as homogeneous.

The samples prepared with the use of hydrated sodium silicate (ester technology), furfuryl resin, and alkdy resin were prepared according to the binder producers’ instructions using a laboratory mixer with a 4-kg capacity and compacted using an LUZ-1 laboratory vibration device. The molding sand used for the samples for cold-box technology was prepared in the same laboratory mixer but was not followed with compaction. The sand was placed in the MULTISERW-Morek Company Universal Core Shooter. The shooting parameters were set as follows: pressure – 0.5 MPa; time – 2 s; amount of amine – 2.47 ml/1 kg of molding sand.

The density of the samples oscillated from 1.48 g/cm³ to 1.63 g/cm³. The sample population varied. For each graph a minimum of three samples that did not differ more than 10% in terms of the obtained results, was taken into account.

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The obtained results are illustrated in Figures 8–10. Test samples showed analogical behavior in the bending strength test (Fig. 8) as they did in the elasticity tests (Fig. 9).

The highest results were achieved for the molding sands prepared in the cold-box technology, reaching 4.23 MPa. This was 69% more than for the molding sands with hydrated sodium silicate (which achieved the lowest value of 1.30 MPa). The molding sand with furfuryl resin reached a bending strength result of 2.71 MPa, and the molding sand with the alkyd resin reached 3.68 MPa (which is, accordingly, 36% and 13% lower than the obtained maximum).

The results of the elasticity measurements give two types of information: the first being the maximum load \( F [N] \) that a sample can withstand, which is analogical to a sample’s bending strength \( R_g [\text{MPa}] \) and maximum deformation \( D_E [\text{mm}] \), and the other is related to the deformation of different samples under a set force value (relating to the way the automated core assembly is realized).

The obtained curves (Fig. 9) visualize a few stages of the elasticity test. First, we can observe a nearly linear propagation of the deformation with increasing force. Second, we can observe a bend in the curve, which is the moment of sample breakage. The short growth of the curve after the maximal strength that the sample can withstand is not taken into the analysis – it is correlated with the eversion of the indenter.

As can be seen in Figure 9, the molding sand with hydrated sodium silicate as the binder achieved both the lowest deformation (0.21 mm) and load resistance (64 N). The molding sand bonded with the furfuryl resin obtained a value of 0.25 mm deformation and withstood 146 N of force. The molding sands bonded with the alkyd resin achieved the result of 0.36 mm in the bend under a force of 181 N. The highest obtained results both in deformation and load size were achieved in the molding sand prepared in the cold-box technology; deformation reached 0.55 mm under a force of 214 N just before the sample’s destruction. The resulting deformation was nearly 62% greater than the molding sands with the hydrated sodium silicate (under more than 70%-greater force).

Based on both elasticity and bending strength, it can be easily deduced that the greater the bending strength and overall strength resistance of the molding sand, the greater deformation the sand will undergo and withstand. However, this is only an analysis of the final outcome after destruction of the samples. What has to be pointed out is the deformation of the samples made from different molding sands under the same force.

If, for example, the impact of a force of 50 N on the deformation of a chosen molding sand would be taken into account, it can be seen that the highest deformation under this pressure is 0.19 mm for molding sands with the hydrated sodium silicate. At the same time, the molding sands with organic-type binders reaches very similar results of 0.11 mm under the same force of 50 N, which is roughly half of what was obtained for the inorganic sand mixture [21].

The last stage of the research presented in this paper is the measurement of molding sand thermal deformation by determining the hot distortion parameter. Thanks to the latest alterations in the measuring equipment, it was possible to mount the sample and pre-heat the equipment without interfering with the sample. This increased the repeatability of the measurement. The lower (bottom) heating plate in the measuring device was pre-heated to 900°C and then placed below the sample. Both the time and temperature were measured on the sample level.

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The results are illustrated in Figure 10.

The thermal deformation of the molding sand with the furfuryl resin has a typical pattern, with an intense deformation of the sample (exceeding 4 mm) and its sudden destruction at a temperature of about 325°C – the deformation was in the opposite direction to the heat source. The deformation of the molding sand with the alkyd resin exceeds approx. 1.5 mm, and the sample collapses at 266°C – the direction of the deformation was headed towards the heat source. The thermal deformation tests (hot distortion parameter) showed that the molding sands with the hydrated sodium silicate and those prepared in the cold-box technology are characterized by better heat stability than the molding sands with the furfuryl and alkyd resins. Both of the tested mixtures exhibit almost no thermal deformation at a temperature range of 0 – approx. 160°C. After crossing the above-mentioned temperature, the samples are subjected to mild deformation until they are damaged. The molding sand prepared in the cold-box technology is characterized by the longer time needed for the sample’s destruction (about 55 seconds), while the sample made from the molding sand with the sodium silicate begins to degenerate after about 15 seconds. This can be advantageous in terms of the time of contact of the molding/core sand to elevated temperatures during and after the pouring process.

4. CONCLUSIONS

Our own research presented in the paper showed the following:

- There are various differences in the behavior of molding sands exposed to bending forces depending on the type of binder used.
- The presented research proves that the change of the used binder and technology influences the level of molding sand deformation at ambient temperatures.
- The conducted research shows that molding sands with inorganic binders that tend to have cohesive type of breakage, achieve lower values in both deformation at ambient temperatures and bending strength when compared to organic binders with the adhesive type of breakage.
- Molding sands with hydrated sodium silicate and molding sands prepared in the cold-box technology are characterized by better heat stability than molding sands with furfuryl and alkyd resins. The longer time needed for the sample destruction achieved for the molding sand prepared in the cold-box technology can be advantageous in terms of the time of contact of the molding/core sand to elevated temperatures during the pouring process.

The presented article shows only a fragment of the broader research on this topic. The need for seeking new testing methods is clearly visible, and the parameter can prove to be a valuable asset in choosing the optimal molding sands for highly mechanized foundries.

Acknowledgements

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