WATER RESISTANCE OF STRUCTURED SAND-SODIUM-SILICATE MIXTURES

**Purpose.** To establish regularities of changes in sand-sodium-silicate mixtures (SSSM) relative water resistance, structured by steam-microwave solidification method (SMS), on their structuring parameters and sodium silicate solute (SSS) dissolution conditions.

**Methodology.** Technical purity water, SSS with 2.8–3.0 silicate modulus, quartz sand with 0.23 mm average particle size were used in this work. Studies were carried out on quartz sand samples, which were preliminarily cladded with 0.5–2.5 % (by weight) SSS and structured by SMS method. The fracture time of structured mixtures was evaluated on cubic specimens with 20 mm rib length, which were immersed in water at different temperatures.

**Findings.** Changes in SSSM water resistance in water regularities depending on their manufacturing, use and storage conditions were established. Recommendations for SSSM rods removal from castings in water were developed.

**Originality.** For the first time, it has been established that relative water resistance of the SSSM structured by SMS method decreases along with increase in water temperature in which it is destroyed. Depending on SSS mass used for quartz sand cladding in range 0.5–2.5 %, water resistance dependence on treatment in steam-microwave environment with 2–7 minutes’ running time acquires an inversion character with inversion point ~3.1 minutes into the processing.

For the first time, it has been discovered that in boiling water the relative water resistance of SSSM, structured by SMS-process and having preheating, monotonically increases with preheating temperature rising from 100 to 600 °C, sharply increases when heated to 600–700 °C and practically becomes absolute after preliminarily preheating to higher temperatures.

**Practical value.** Research results will be useful in concepts of processes accompanying destruction of structured sands with water-soluble binders expanding, as well as in technologies and equipment development designed for SSS rods and mold rests removing from castings.

**Keywords:** water resistance, sodium silicate solute, sand-sodium-silicate mixture, knockout, steam-microwave solidification, rod binding materials, are environmentally and sanitary-hygenically safe. Currently, in foundries, share of SSS used in CMC production is not large. This is mainly due to the high energy consumption and, in some cases, impossibility of sand-sodium-silicate mixture (SSSM) removing from castings [8, 9]. Therefore, most of foundries for CMC manufacturing use synthetic resins, which, with all their environmental and economic disadvantages, make it possible to obtain CMC with a higher level of technological properties. Using resin in CMC is accompanied by a large amount of carcinogenic and toxic gaseous substances (formaldehyde, phenol, benzene, and so on) polluting into foundries working areas and, accordingly, into the environment. At the same time, release of such substances into air begins from the moment of CMC manufacturing, increases sharply from the moment of pouring to the moment of knocking out the casting from mold and rod and does not finish even after disposing spent mixture in dumps [10]. To solve this environmental problem, instead of organic origin binding materials, a number of authors now recommend returning to use inorganic binding materials, in particular, to phosphate binders, including aluminum-chromium phosphate, aluminum-iron phosphate, aluminum-boron phosphate concentrate with powder-solidifier based on magnesium oxide, phosphoric acid with chemically active dust addition to CMC, and so on or SSS. Possessing obvious environmental advantages over organic binders, phosphate-sand mixtures, however, are characterized by unstable properties and acidic properties undesirable for use, regeneration and disposal of waste mixtures. Sand, clay and water mixtures are structured in conductors by cooling them down to 15–75 °C. This creates danger of frostbite for molds and requires expe-
sive refrigeration equipment. Mixtures with gypsum and cement also have a number of their own characteristics and conditions of implementation, are characterized by significant dust emission during their preparation and knocking out castings from them, as well as laboriousness of regeneration. That is, from this point of view SSSM is more promising.

SSSM structuring is carried out after their manual or mechanical compaction in equipment by heat drying (dehydroation), blowing with carbon dioxide, due to adding powdered dicalcium silicate, complex ether, and so on to the mixture. The main disadvantages of such mixtures are major efforts of castings mechanical knocking out and/or sand regeneration laboriousness. In this regard, wasted SSSMs are disposed without regeneration at landfills, in dumps, are partially used in road construction, and so on.

Summarizing results of numerous publications and studies, Lucarz M. (2008) in his works notes that disposal of waste CMC in dumps or its use in road construction, to fill closed mines cavities, etc. poses critical threat for environmental safety. Therefore, waste quartz sand regeneration is a mandatory technological operation that should be performed not only for its recycling, but also in terms of its subsequent safe disposal.

In order to facilitate SSSM knocking out from castings, it is proposed to add certain technological additives to SSS, modify SSS during its preparation, reduce SSS content in mixture, and so on [7, 8, etc.]. At the same time, the authors of these works consider SSSM removal from castings solely from viewpoint of shock and vibration effects on them.

In accordance with data [14], SSSM knocking out work (A) depends on method for its structuring, heating temperature and can be described by dependencies whose characteristics forms are presented in Fig. 1.

According to authors [14, 15] (Fig. 1), maximum upon SSSM heating to 200 °C is consequence of sodium silicate and SiO2 gel mixture in cuffs formation. SSSM heating to 600 °C leads to de-strengthening and gel destruction as a result of SSSM thermal expansion. When heated to 800 °C, SSSM knocking out work increases again, which is due to sodium silicate melting, formation of more perfect-shaped cuffs and their solidification upon subsequent cooling. Mixtures heating to 900–1100 °C and their subsequent cooling are accompanied by partial silicon dioxide dissolution from sand grains in molten sodium silicate and its subsequent devitrification (partial crystallization), which leads to SSSM knocking out work decreasing, when heated to these temperatures.

To reduce knocking out work, many scientists and researchers recommend reducing water glass content in mixture to 2–3 % (by mass), using technological additives in SSSM (carbon-containing materials, metallurgical dust, sulphite waste liquor, wood and cardboard dust, iron sulfate hydrate, kaolin, and others). Authors [8] recommend using exclusively pore-forming or thermally expanding materials as such additives for SSSM.

A large number and variety of recommended technological aids in SSSM are mainly due to:

- peculiarity of SSS and sodium silicate physicochemical properties;
- technological limitations and peculiarities of using technological additives;
- instability of properties and efficiency of technological additives from natural and technogenic origin materials;
- expensiveness of industrially produced technological additives.

The disadvantages of using any technological additives include:

- necessity of energy-intensive sand regeneration for its recycling;
- lack of comprehensive solution to problems of low manufacturability, inherent in sand-sodium-silicate CMC.

That is, no technological additive in SSSM will fully provide a required level of processability for mixtures and CMC made from them.

SSSM structuring by SMS-process is one of ways of comprehensive solution to this problem. Since when used in mixture of less than 2–3 % (by weight) SSS without any technological additives in SMS provides opportunities of:

- filling the rigging with mixture by free pouring and vibration compaction;
- fast mixture in rigging structuring (solidifying);
- exclusion of CMC drying process;
- process of knocking out castings from castings molds and cores low energy consumption;
- multiply reusing of waste mixture after grinding and sieving in CMC production;
- elimination of necessity for regeneration and disposal of spent mixture due to its use as raw material for subsequent silicate block production.

Unlike dependences 1–3 in Fig. 1, dependence 4 has no maxima and testifies to the low value of SSSM structured by microwave radiation knocking out work when it is heated to temperatures above 300 °C. M. Stachowicz, K. Granat, D. Nawak (2011) see such difference in nature of course dependences in Fig. 1 in structure peculiarities of sodium silicate cuffs between sand grains, which are formed under microwave radiation on the mixture action.

Depending on mass ratio, wall sizes, pouring conditions, melt poured into the mold temperature and a number of other parameters, the temperature of the rod’s various parts can vary from the initial rod temperature before melt pouring into the mold moment to the liquidus temperature of alloy poured into the mold. Based on this, a number of areas can be distinguished in sand-sodium-silicate rods, whose diagrams are shown in Figs. 2, a–f, and character of temperature distribution over the cross section of the rod for corresponding schemes is shown in Fig. 2, e.

If it is assumed that temperature td is heating temperature at which subsequently cooled sodium silicate in cuffs of structured mixture loses its ability to dissolve in water within technologically acceptable time, then it follows that sand-sodium-silicate rods structured as result of sodium silicate dehydration, under certain conditions, can be completely (Fig. 2, a) or partially (Fig. 2, b) removed from castings not only by shock-vibration action on castings, but also as a result of their contact with water. In all other cases (Figs. 2, c–f), rods removal should be carried out by shock-vibration action on castings.

Ability to remove rods by immersing them in water is important not only for castings from alloys with low hardness (alloys based on copper, aluminum, zinc, lead, and others), but also with low ductility (cast iron), etc. Nevertheless, until now, there are no data on possibility of removing sand-sodium-silicate rods structured in steam-microwave environment from castings. This reduces commercial attractiveness of this method for producing rods and does not contribute to solving the problem of lowering energy consumption of technological operation of knocking out sand-sodium-silicate rods from castings, generally.

In this regard, search for new technical solutions to reduce energy consumption of technological operation of knocking out sand-sodium-silicate rods from castings, in particular, by
processing them in water, is an urgent problem of scientific and applied importance.

**Literature review.** SSS specificity as a binder material is its solidification possibility, both as a dehydration result [15] and as a result of chemical reactions leading to sharp increase in its silicate modulus [16, 17]. It follows from this that sand–sodium-silicate rods, structured by their dehydration, under certain conditions, can be removed as a result of their immersion in water.

According to V. I. Korneev, V. V. Danilov (1996) SSS with silicate modulus of 2.8–3.0 is sodium silicate water solution, which, in addition to oligomers, contains silica with high degree of polymerization and high cohesion. That is, it is characterized by certain ratio of molecular (monomeric – α-SiO₂) and colloidal (β-SiO₂) silica. At the same time, presence of sodium dioxides in dry SSS provides not only its dissolution possibility in water, but also control of this process rate.

According to Waldes H. H. and Lange K. R. (1969), dissolution rate of solid SSS particles in water increases with SSS silicate modulus decreasing, with water temperature increasing, with presence of hydrated water (18.5 % by weight) in dry SSS, with dissolved SSS particle size decreasing. Presence of hydrated water in sodium silicate molecules composition is fundamental condition for its rapid dissolution in water [18]. Thus, it will be assumed that loss of SSS solubility occurs with its complete dehydration. Complete dehydration of SSS, according to V. I. Korneev and V. V. Danilov (1996), occurs when SSS reaches 600 °C. Then, SSS should not dissolve in structured SSSM, even briefly heated to temperatures above 600 °C. Accordingly, sand-sodium-silicate rods structured by SSS dehydration and not heated to more than 600 °C should be removed from castings as a result of their immersion in water.

Korneev V. I. and Danilov V. V.’s (1996) research results showed that SSS dissolution rate can be significantly influenced by surface carbonization of SSS particles dissolved in water. Carbonation is a process whose activity increases after SSS loses its free water and continues until moment when hydration water from sodium silicate removes completely [19]. That is, gradual loss of hydrated water by amorphous silicate solutions with silicate modulus of about 3, solidified in air, is accompanied by carbon dioxide SSS absorption from air and by appearance of chemical compounds NaHCO₃, Na₂CO₃. This process leads not only to sodium ions migration from deep regions to particles surface and crystalline carbonate structures appearance, but also to low–water silica framework formation, as a result [20]. Such framework appearance increases system under consideration water resistance and, therefore, decreases solid SSS particles in water dissolution rate. It follows from this that increase in duration of rods holding in air should lead to their water resistance and, accordingly, increasing duration of removal of rods from castings in water.

It follows from above that water resistance of SSSM structured by dehydration depends on a large number of factors, including method of their solidification. At present, there are no data on SSSM structured in steam-microwave environment (method of steam-microwave solidification – SMS) water resistance and effect of structuring conditions on their water resistance.

**Methods.** Technical purity water, sodium silicate solute with silicate modulus of 2.8–3.0 and specific density of 1.44 g/cm³, quartz sand with an average particle size of 0.23 mm and clay content of up to 0.2 % (by mass) were used in this work.

Mass was determined on electronic balance with an accuracy of 1.0, 0.1 and 0.01 g. Dimensions were measured with Vernier caliper with an accuracy of 0.1 mm. Time was recorded with stopwatch with an accuracy of 1 s.

Studies were carried out on cubic specimens with rib length of 20 mm. Test samples were obtained by cutting blank weighing 2.2 kg from mixture structured by SMS-process. For blank, quartz sand cladded with sodium silicate solute 0.5–2.5 % (over 100 % sand) was used. Cladded sand was dried in air to water mass content in sodium silicate of 17–19 %. After air drying, sand was dried additionally for 5 minutes in microwave furnace with magnetron power of 900 W and radiation frequency of 2.45 GHz.

Cladded sand structuring was carried out after sifting it through a sieve with mesh size of 0.4 mm. For structuring, water portion weighing 1 g (water charge – WC) was used, which was placed in foam-polyurethane sponge with dimensions of $230 \times 8$ mm. That is, for structuring, foam-polyurethane sponge filled with 1 g of water was placed on the bottom of a polypropylene box. The box was filled with 2.2 kg of cladded sand and compacted for 30 minutes by vibration with oscillation frequency of 50 Hz and amplitude of 1 mm. After compacting, the box was placed in working space of microwave furnace in the center of a rotating table. While furnace table was rotating at frequency of 2.5 rpm, the box content was treated with microwave radiation for 8 minutes.

To determine destruction duration, testing specimens were installed on X-type support-bracket and in this state have been immersed in water at appropriate temperature. The scheme for cutting cubes from blank samples and the scheme for cube placing in X-type support-bracket are shown in Fig. 3.

Storage of samples before testing was carried out in open air with relative humidity of 60–62 % at temperature of 20–26 °C. Destruction of testing samples was carried out in water with temperature during tests which was maintained at a given value (t) with an accuracy of ±1 °C. Ratio of water mass to tested sample mass in it in all cases was $<100$. For sample destruction duration in water, time of its complete removal from X-type support-bracket was taken.

Samples were preheated to specified temperatures in laboratory resistance muffle furnace according to the following regime:
- rate of temperature rise in furnace working space – 80–170 °C/h;
- isothermal holding duration at given temperature – 40 ± 1 minutes;
- cooling in air outside the furnace from predetermined temperature until the sample reaches 20–22 °C. Error in temperature determining during samples isothermal holding was less than 3 °C.

Dependence of hydrated water content in structured SSSM sodium silicate cuffs (WC weight during SSSM structuring was 0.5 g) was estimated from results of SSSM samples with mass 200–230 g weighing, heated in microwave furnace for given time. At the same time, duration of steam-microwave treatment (t) was taken into account, with SSS mass used for cladding quartz sand of 0.5 and 2.5 % (over 100 % of the sand). Hydrated water mass in sodium silicate SSSM after microwave treatment was determined by the sample treated with microwave radiation weighing before and after heating it to 700 °C in resistance furnace, with following it recalculation to dehydrated sodium silicate mass in test sample.

**Results.** In accordance with research methodology, relative duration value of samples destruction in water (water resistance) was calculated by formula

\[ V = \frac{\tau_i}{\tau_i} \]

where \( \tau_i \) is destruction duration of the \( i^{th} \) sample in water, \( \tau \) is destruction duration of samples in water (water temperature 80 °C) from quartz sand structured by microwave radiation for 2 minutes (\( \tau = 94 \) s), cladded with 0.5 % (by mass) SSS.

Relative dependence of relative destruction duration in boiling water (water resistance) of samples with 1.5 % SSS, structured by SMS-process for 2, 4, and 7 minutes, on their temperature preheating is shown in Fig. 4.

From analysis of dependence course in Fig. 4 it follows that with preheating temperature increasing from 100 to 600 °C, relative duration of samples destruction (water resistance) monotonically increases and can be calculated by formula

\[ V = 1.32 \cdot \ln (t) - 5.69 \]

where \( t \) is sample preheating temperature from 100 to 600 °C.

In preliminary samples heating from 600 to 700 °C case, relative duration of their dissolution (water resistance) increases sharply and when heated to temperatures above 700 °C, samples under considered conditions acquire almost absolute water resistance. In this case, relative duration of samples destruction practically does not depend on their structuring duration according to SMS-process within processing range from 2 to 7 minutes.

This dependence of relative duration of samples destruction on their preheating temperature in range from 600 to 700 °C is explained by dehydration completion and structure of sodium silicate in cubs between sand grains changing.

This conclusion is confirmed by Korneev V. I. and Danilov V. V.’s data (1996), according to which, after heating sodium tri-silicate to 600 °C and its subsequent cooling to room temperature sodium tri-silicate exists in form \( \beta \text{Na}_2\text{O} \cdot 2\text{SiO}_2 + \text{SiO}_2 \), which contains hydroxyl groups. Apparently, exactly this circumstance is the reason for relatively rapid dissolution of \( \beta \text{Na}_2\text{O} \cdot 2\text{SiO}_2 + \text{SiO}_2 \) in water (low water resistance). Upon reaching 600 °C, tri-silicate is almost completely dehydrated and loses its ability to rapidly dissolve in water. At temperatures above 665 °C \( \beta \text{Na}_2\text{O} \cdot 2\text{SiO}_2 + \text{SiO}_2 \) sodium tri-silicate turns into high-silica silicate \( 3\text{(Na}_2\text{O} \cdot 2\text{SiO}_2) + 2\text{SiO}_2 \), which at room temperature loses its ability to dissolve even in boiling water at atmospheric pressure, or it takes a long time to dissolve, or requires increased temperature and pressure of saturated water steam above it. Thus, maximum permissible heating temperature for SSSM, structured according to SMS-process, which ensures its destruction in water in technologically acceptable time, should be considered as: \( t = 600 \) °C. Certain discrepancy between temperatures indicated above and those determined from dependence in Fig. 5 is explained by difference in research conditions and silicate modulus values used in SSS experiments.

Dependence of relative duration of destruction of mixture structured by SMS-process for 2 minutes in water on structured mixture with SSS 0.5 and 2.5 % exposure (storing) duration in air and water temperature at 2 minutes mixture processing in steam-microwave environment, is shown in Fig. 5.

From dependencies in Fig. 5 it follows that relative duration of samples destruction in water (water resistance) increases with decrease in SSS mass used for cladding quartz sand, with decreasing duration of samples exposure in air.

Relative time of dissolution (relative to dissolution time of mixture in water with temperature of 40 to 80 °C after 3–168 hours from the moment of the end of mixture within 2 minutes by the SMS-process structuring and without being heated (Fig. 5) can be calculated using empirical formula

\[ V_2 = \frac{240 \cdot 0.002 \cdot \tau}{(0.1 \cdot \tau_m)^3} \cdot m \]
where \( t \) is structured mixture in air storing duration, hours; \( t_{PR} \) is water temperature during dissolution, °C; \( m \) is SSS mass content in mixture, which was used for quartz sand cladding, % (over 100 % of the sand).

Dependences of relative duration of destruction of structured mixture by SMS-process for 2 and 5 minutes in water on its temperature and the amount of SSS used for quartz sand cladding during initial mixture in steam-microwave environment processing, are shown in Fig. 6.

From analysis of course of the curves in Fig. 6 it follows that with water temperature increasing samples destruction duration (water resistance) decreases. It corresponds to known regularities of influence of solvent temperature on dissolved substance dissolution rate. In this case, if duration of mixture structuring with microwave radiation is 2 minutes, then mixture dissolution duration increases with decreasing SSS mass content, which was used for cladding quartz sand. If mixture structuring duration is 5 minutes, then mixture disintegration duration decreases with SSS mass content decreasing. That is, cladded sand with 2.5 % SSS with SMS-process treatment for 5 minutes was sufficient for residual hydration water in sodium silicate content decreasing to value comparable to amount of residual water in sand cladded with 0.5 % SSS cuffs. It follows from this that with certain duration of SMS-process, the nature of dependences on SMS-process duration and SSS used for sand cladding content should change to opposite.

These regularities are explained by simultaneous influence of two factors on water resistance (V) value — SSS amount used for quartz sand cladding and SMS-process duration during the period of its structuring. To explain these effects let us consider the cuffs schemes shown in Fig. 7.

Distinctive feature of the diagram in Fig. 7, a compared to the diagram in Fig. 7, b is cuff cross-section size. Naturally, with increase in SSS mass, spent on sand cladding, cuff cross section increases. At the same time, limited microwave radiation exposure duration during cladded sand structuring leads to the fact that part of hydration water remains in cuffs with a large cross section. This, in particular, is evidenced by dependence course shown in Fig. 8.

From Fig. 8 it follows that after 2 minutes of microwave treatment in structured sand cladded with 0.5 % SSS cuffs, water content decreases to \( \sim 0.4 \% \). During the same time, in structured sand cladded with 2.5 % SSS cuffs water content reduces to \( \sim 11 \% \). It can be assumed that regardless of initial size, cuffs with bigger amount of hydrated water will dissolve in water at a faster rate than more dehydrated cuffs.

This assumption is confirmed by dependences of relative duration of destruction of structured mixture in water on initial mixture treatment according to SMS-process duration and SSS amount used for quartz sand cladding at water temperatures of 40 and 80 °C, shown in Fig. 9.

From analysis of dependence course in Fig. 9, it follows that, regardless of water temperature in which samples have been destroyed, nature of destruction relative duration (water resistance) from SSS used for sand cladding mass, changes with processing by SMS-process duration \( \sim 3.1 \) minutes. That is, obtained regularities indirectly confirm the assumption about significant effect of structuring duration and, consequently, hydrated water content in SSS cuffs on their water resistance.

**Conclusions.** Sand-sodium-silicate mixture structured by SMS-process water resistance decreases with water temperature increasing and acquires an inversion character from its duration within 2–7 minutes and sodium silicate solute mass used for quartz sand cladding within 0.5–2.5 % (over 100 % of the sand). Inversion point corresponds to \( \sim 3.1 \) minutes of SMS-process duration. These peculiarities are due to residual (hydration) water amount in sodium silicate, which is located in structured mixture cuffs, size of cuffs themselves, which, in turn, depend on SSS consumed for cladding quartz sand mass.

With increasing duration of storing sand-sodium-silicate mixture structured by SMS-process in air, its water resistance increases, which is due to hydrated sodium silicate carbonization process in cuffs between grains of structured mixture.

Water resistance of sand-sodium-silicate mixture, structured according to SMS-process and preheated from 100 to 600 °C, in boiling water monotonically increases with heating temperature increasing, sharply increases at temperatures from 600 to 700 °C and becomes almost absolute at higher temperatures.
Водостійкість структурованих піщано-рідкоскляних сумішей

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Мета. Встановити закономірності зміни відносної водостійкості піщано-рідкоскляних сумішей (ПРС), структурованих методом паро-мікрохвильового затвердження (ПМЗ), від параметрів їх структурування та умов розчинення рідкого скла (РС).

Методика. У роботі використовували воду технічної чистоти, РС із силикатним модулем 2,8–3,0, кварцовий пісок із середнім розміром частинок 0,23 мм. Дослідження проводили на зразках із квартового піску, що попередньо був плаціваної 0,5–2,5 % за масою, РС і структурований методом ПМЗ. Час руйнування структурованих сумішей розраховували на зразках кубічної форми з довжиною ребра 20 мм, зафіксованих у воді за різних температур.

Результати. Встановлені закономірності зміни водостійкості ПРС у воді в залежності від умов їх виготовлення, використання та зберігання. Розроблені рекомендації з використання та зберігання. Розроблені рекомендації з видалення піщано-рідкоскляних стрічок iз виливок у воді.

Наукова новизна. Уперше встановлено, що відносна водостійкість ПРС, що структурована методом ПМЗ, зменшується зі збільшенням температури води, в якій вона руйнується. У той же час, у залежності від маси РС, внаслідок для плацівання кварцового піску в діапазоні зони від 0,5 до 2,5 %, залежність водостійкості від обробки паро-мікрохвильового середовища тривалістю від 2 до 7 хвилин набуває інверсійного характер з точкою інверсії на ~3,1 хвилини обробки. Уперше встановлено, що в кипячій воді у ПРС, що була структурована за ПМЗ-процесом і прийшла попередній нагрів, відносна водостійкість монотонно зростає при підвищенні температури попереднього нагрівання від 100 до 600 °С, а при нагріванні до 600–700 °С різко зростає i практично стає абсолютною після попереднього нагрівання до близько високих температур.

Практична значимість. Результати досліджень будуть корисні в частині зもなく зі збільшенням температури води, в якій вона руйнується. У той же час, у залежності від маси РС, збільшення вологоності паро-мікрохвильового середовища тривалістю від 2 до 7 хвилин набуває інверсійного характер з точкою інверсії на ~3,1 хвилини обробки. Уперше встановлено, що в кипячій воді у ПРС, що була структурована за ПМЗ-процесом і прийшла попередній нагрів, відносна водостійкість індукована підвищенням температури попереднього нагрівання від 100 до 600 °С, а при нагріванні до 600–700 °С різко зростає і практично стає абсолютною після попереднього нагрівання до близько високих температур.

Ключові слова: водостійкість, рідків склі, піщано-рідкоскляна суміш, вибраківка, паро-мікрохвильове затвердження, стрічка

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