# Polymeric superplasticizers based on polycarboxylates for ready-mixed concrete: current state of the art

#### Daniel Wiliński<sup>1), \*)</sup>, Paweł Łukowski<sup>1)</sup>, Gabriel Rokicki<sup>2)</sup>

DOI: dx.doi.org/10.14314/polimery.2016.474

**Abstract:** Chemical admixtures are one of the basic components of the contemporary concrete. The most important admixtures are superplasticizers, *i.e.* the modifiers which allow for high-range reducing of the water content in the concrete mix. The most effective superplasticizers are based on polycarboxylates and their derivatives. The state of the art in the range of polycarboxylate admixtures for concrete is presented in the paper. The methods of synthesis and modification of carboxylic polymers are characterized and the mechanism of action of polycarboxylate superplasticizers is described. The influence of the chemical structure of polycarboxylates on the consistence of the concrete mix, as well as the prognosis of the superplasticizers development is also presented.

Keywords: superplasticizers, polycarboxylate, chemical admixtures, cement, concrete.

## Polikarboksylanowe domieszki upłynniające mieszankę betonową: aktualny stan wiedzy

**Streszczenie:** Podstawowym składnikiem betonu są obecnie domieszki chemiczne. Za najważniejsze uznaje się superplastyfikatory, czyli domieszki silnie redukujące zawartość wody w mieszance betonowej. Do najefektywniejszych znanych superplastyfikatorów zaliczają się polikarboksylany i ich pochodne. Artykuł stanowi przegląd aktualnej literatury dotyczącej polikarboksylanowych domieszek chemicznych. Scharakteryzowano metody syntezy i sposoby modyfikacji polimerów z grupami karboksylowymi i omówiono mechanizm działania takich superplastyfikatorów. Przedstawiono zależność między strukturą chemiczną polikarboksylanów a stopniem upłynnienia mieszanki betonowej oraz prognozę rozwoju superplastyfikatorów.

Słowa kluczowe: superplastyfikatory, polikarboksylany, domieszki, cement, beton.

In the current concrete technology, the chemical admixtures are important components of the concrete mix. The main progress in the concrete technology is associated with the admixtures development. The admixture for concrete is defined as the material added to the concrete mix in a quantity not more than 5 wt % of the cement mass. Therefore, the concentration of the polymer is too low to form the film inside the hardening cement paste; this can take place when the content of polymer is higher (in such case the polymer modifier is called an addition) [1]. Most chemical admixtures applied in concrete present anionic polyelectrolytes which physically adsorb on the cement grain surface for effective acting. Such admixtures aimed at modification of the concrete mix properties were first introduced in thirties of the last century. Their function was to improve the consistence of the cement paste [2]. Figure 1 clearly indicates that the introduction of superplasticizers, which made possible a decrease of water to cement ratio (w/c), afforded a significant increase in the concrete strength [3]. Nowadays, the use of admixtures varies greatly from one country to another. For example, in Japan almost 100 % of concrete contains admixtures. But, for example, in the United States more than 50 % of concrete contains chemical agents, although this percentage is increasing constantly as the beneficial role of admixtures is understood [4]. Apart from concrete, polycarboxylate (PCE) superplasticizers are also used in the manufacture of gypsum materials [5].

The admixtures affecting the consistence of the concrete mix can be categorized as water reducing admixtures (plasticizers) and high-range water reducing admixtures (superplasticizers, also called the liquefying admixtures), the latter being much more effective. The efficiency of the admixtures is evaluated by the possible

<sup>&</sup>lt;sup>1)</sup> Warsaw University of Technology, Faculty of Civil Engineering, Department of Building Materials Engineering, Al. Armii Ludowej 16, 00-637 Warsaw, Poland.

<sup>&</sup>lt;sup>2)</sup> Warsaw University of Technology, Faculty of Chemistry, Noakowskiego 3, 00-664 Warsaw, Poland.

<sup>\*)</sup> Author for correspondence; e-mail: d.wilinski@il.pw.edu.pl

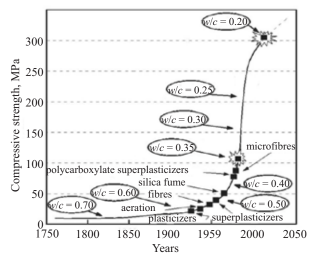


Fig. 1. Generalized curve of concrete development [3]

reduction of the water content in the concrete mix, with keeping the same consistence. This reduction (as compared to the control mix) should be  $\geq 5$  wt % for the plasticizers and  $\geq 12$  wt % for the superplasticizers [6]. The main goals in the use of the superplasticizers are:

- improvement of the concrete strength (due to the reduction of the water content),

– improvement of the concrete mix consistence without change of the concrete composition (w/c = const),

– decrease in the cement consumption without affecting the strength of the concrete and consistence of the concrete mix.

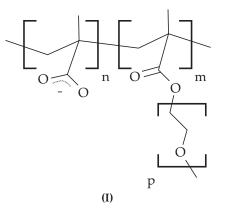
The effectiveness of superplasticizers depends on various technological factors. These factors could be divided into three groups:

 – concrete constituent characteristics (mainly cement, superplasticizers and mineral additives characteristics),

 mixture proportions (water to binder mass ratio, superplasticizer, cement and mineral additives content, presence of other chemical admixtures in mixture),

– the methods and conditions of concrete production (order of addition, temperature of the mixture) [7].

The polycarboxylate admixtures, based mainly on the polymers with carboxylate and oligo(ethylene oxide) side groups, are one of the most effective superplasticizers from among all known modifiers; they are usually referred to as the liquefying admixtures of a new generation. The



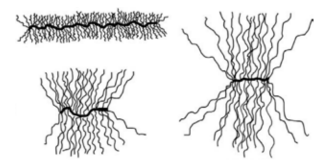


Fig. 2. Schematic examples of molecular architecture of PCE superplasticizers [8]

methacrylate copolymers of methacrylic acid and methacrylic ester with the oligo(ethylene oxide) group is an example of such admixture [Formula (I)], commonly called the polycarboxylate (PCE) superplasticizer. It is a comblike polymer built up with ionic hydrophilic carboxylate groups (COO<sup>-</sup>) and long, elastic side nonionic hydrophilic oligo(ethylene oxide) chains Fig. 2 [8].

Superplasticizers are useful in obtaining a dispersion of the cement grains in water. The carboxylate groups (COO<sup>-</sup>) interact with the surface of the cement grain, which leads to polymer adsorption and electrostatic repulsion between grains (COO<sup>-</sup> groups create a negative charge around the grain). In many publications, also a steric repulsion is considered as a potential mechanism of cement grains dispersion. Detailed information about interaction mechanism of cement and superplasticizers can be found in [5, 9, 10]. Also, the wettability of grains is improved, fostering the full cement hydration (setting). The viscosity of the concrete mix decreases leading to the consistence improvement. The research by Borget et al. [11] showed behavior of PCE superplasticizers (PCEs) in different salt solutions. In classical cement--based media at room temperature, PCEs behave generally as polymers in good solvent, except in case of very high sulfate concentration. Duan et al. [12] studied the molecular polarization of polycarboxylate by measurement of electrical conductivity in deionized water solution. It was found that the PCE significantly increased the conductivity of the solution with maximum values reached at PCE concentrations between 20 and 25 wt %. These results indicate that the degree of polarization was in connection with the molar mass of the polymers.

Numerous studies, for instance Winnefeld *et al.* [13], Peng et al. [14], Kjeldsen *et al.* [15], Ran *et al.* [16], Zingg *et al.* [17, 18], Rai and Gajbhiye [19], Üzer and Plank [20], have proved that the superplasticizers in the cement pastes directly influence their rheological properties. It is commonly accepted that the influence of superplasticizers on rheology of cement binder mixture depends primarily on the superplasticizer and cement characteristics and their interaction. Effects of PCEs on increasing workability [13, 21, 22] and fluidity [14, 23–33] of fresh concrete have already attracted many attentions so far. The authors reported that PCEs have a good performance on

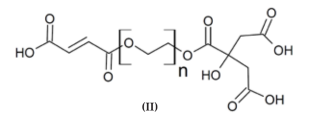
Macromonomer introducing			
oligo(ethylene oxide) side cha- ins into polymer	Monomer introducing carboxyl groups into polymer	Reference	
	он	[26, 29, 47, 53]	
	-он	[12, 16, 21, 37, 39, 54]	
≈ llr ~ lo.	-он	[21, 38]	
	-он	[27]	
"ДС. Лон		[19, 48, 55]	
1	-он	[21, 33, 42, 56]	
↓[ <sup>0</sup> →]n	он	[9, 11, 13–15, 17, 18, 20–23, 36, 40, 45, 48, 49, 53, 57–66]	
	он	[12, 28]	
		[67]	
≈ ~T ~ ToH	$\sim \sim $	[24, 25, 30, 68]	
	но	[30]	
	-он	[21, 39]	
محرق∽]₀́	-он	[21]	
	°Z°F°	[8, 21, 35, 46, 59, 64, 65]	
<sup>↓</sup> [° → ] <sub>°</sub>	-он	[27, 31]	
	Сн	[21, 44]	
↓[°~},~	-у-он	[21, 44]	
	°₹°≻°	[21, 60]	
	——————————————————————————————————————	[32, 69]	
л с с Тон И	$\sim \sim $	[32]	

### T a b l e 1. Monomers for synthesis of polycarboxylate-type superplasticizers with oligo(ethylene oxide) chains

#### Table 1. (Continued)

Macromonomer introducing oligo(ethylene oxide) side cha- ins into polymer	Monomer introducing carboxyl groups into polymer	Reference	
<pre>&gt;&gt;_[o→]_oH</pre>	-он	[31]	
Ŷ~-[°~]₀~	ОН	[43]	
√[°√] <sub>OH</sub>	Сн	[12]	
	он	[23]	
	°₹°≻°	[57]	
	-он	[41]	
≈.√Ìţo.\]"	он	[41]	

improving workability of concrete and fluidize the mixture more effectively than other superplasticizers. Some types of superplasticizers cause the rise of excessive air content [34] and exhibit foaming action [35]. Morin et al. [36] and Chen [37] analyzed the influence of PCEs on the shrinkage of the concrete. The shrinkage partly compensates the volume augmentation during mixing caused by the air entrapment induced by the superplasticizer. There are many works [9, 33, 38-44] devoted to effects of polycarboxylate superplasticizers on the hydration behavior of cement paste, however, influence of molecular structure of PCEs on the hydration behavior of cement remain poorly understood. In addition to influence of PCEs on hydration of cement, effects of PCEs on microstructure of cement paste and durability of concrete were reported as well [17, 30, 45, 46]. The research by Huang et al. [46] showed that the mortars with PCEs have lower porosity than that with conventional superplasticizers based on naphthalene (PNS). The fraction of pores with diameters larger than 100 nm in the mortars with PCEs is lower than in the mortar with PNS. It can be concluded that PCEs leads to denser microstructure in concrete than PNS and, therefore, better durability performances of concrete. The compressive strength of concrete, influenced by PCEs was investigated by Li et al. [32] and Wu et al. [47]. They have found that compressibility of the concrete mixture increased after adding PCEs (after 28 days reached even 30 % more compressive strength than concrete without PCEs). According to Winnefeld et al. [13], Zingg et al. [18] and Plank et al. [48] the presence of PCEs not only shift the main peak of heat of hydration of cement pastes but also changes the peak shape.



#### METHODS OF POLYCARBOXYLATE SYNTHESIS

The most commonly used side chain for PCE superplasticizers is oligo(ethylene oxide). The polymers with carboxylate and oligo(ethylene oxide) groups can be obtain by radical copolymerization of monomer introducing the carboxylic group into polymer (*e.g.* acrylic acid) with macromonomer introducing the oligo(ethylene oxide) chain into polymer [*e.g.* oligo(ethylene oxide) acrylate].

The optimal esterification condition of macromonomer has been studied by Jiang *et al.* [49]. The monomers used for manufacturing of the polycarboxylate superplasticizers are collected in the Table 1.

It should be mentioned that there are other ways of PCE synthesis. Lu *et al.* [26] obtained polycarboxylate superplasticizers by terpolymerization of macromonomer – monoester of maleic acid [Formula (II)], oligo(ethylene oxide) acrylate and acrylic acid.

Liu *et al.* [50] used poly(acrylic acid) and poly(ethylene oxide) terminated with amino and methoxy groups as reactants for synthesis of amide-PCE (Scheme A).

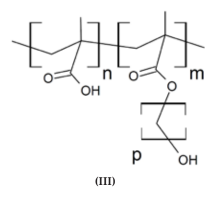
Lei and Plank [51] reported on a structurally modified PCE superplasticizer which was synthesized from methacrylic acid and 2-hydroxyethyl methacrylate, 3-hydroxypropyl methacrylate or 4-hydroxybutyl methacrylate esters. These PCEs possess hydroxyalkyl side chains [Formula (III)].

In other papers Lei and Plank presented synthesis of terpolymers from maleic anhydride, monoalkyl maleate esters, and 4-hydroxybutyl vinyl ether [52]. Taking into account the above considerations, we propose to classify the PCEs as the following five categories:

– **EsMa** – type PCEs, when **Es**ters and **M**onoprotic **a**cids are used as substrates,

– **EsPa** – type PCEs, when **Es**ters and **P**olyprotic **a**cids or anhydrides are used as substrates,

– AlMa – type PCEs, when Allyl or ether compounds and Monoprotic acids are used as substrates,



– AlPa – type PCEs, when Allyl or ether compounds and Polyprotic acids or anhydrides are used as substrates,

- Amide - type PCEs, when amide bonds are formed.

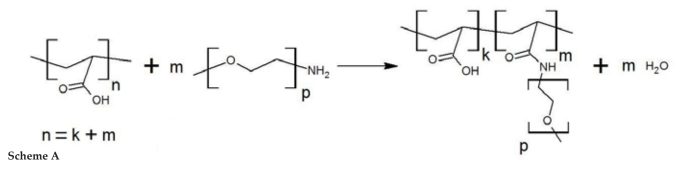
Table 2 presents examples of each type of polycarboxylates. The most commonly used PCE superplasticizers are classified as EsMa-type.

#### MODIFICATION OF POLYCARBOXYLATES

At present, polycarboxylate superplasticizers are modified by insertion of other monomers to afford terpolymers (Table 3). The particular attention is also paid to versatility of the liquefying admixtures, *i.e.* their high effectiveness with various types of cements. For instance, the polycarboxylates modified with methoxysilanes [Formula (IV)], contrary to the traditional superplasticizers, are very effective when the cements with high sulphates content are used [22]. In the field of waterreducing admixtures, the research and development of newer polymers is being undertaken utilizing tailored superplasticizers [70].

#### INFLUENCE OF THE CHEMICAL STRUCTURE OF POLYCARBOXYLATES ON THE LIQUEFYING PROPERTIES OF CONCRETE MIX

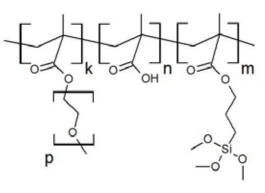
The molecular structure of PCE have a close relationship with their performance in cement system. Peng *et al.* [61] showed that the effectiveness of the polycarboxylate superplasticizers grafted with poly(ethylene oxide) depends strictly on the molar mass of the polymer used. In other research, Peng *et al.* [14] synthesized six polycarboxylates with similar molar mass and dispersion



e polycarboxylate superplasticizers Type of PCEs Example					
EsMa	Example				
EsPa	HO OH C				
AlMa					
AlPa					
Amide					

#### T a b l e 2. Examples of EsMa, EsPa, AlMa, AlPa and amidetype polycarboxylate superplasticizers

#### and constant length of side chains of poly(ethylene oxide) [p = 45, Formula (I)], varying in the content of the carboxylate groups (*i.e.*, molar ratio of the monomers used) (Table 4). The consistence of the cement mortar *vs*. amount of the admixtures as well as ability of the particular superplasticizers to reduction of the water content in the mix are presented in Fig. 3. The optimum molar ratio of the monomers n/m in PCE was 3.6.



T a b l e 3. Monomers used as modifiers for polycarboxylate superplasticizers				
	Monomer			
Compound name	Chemical structure	Reference		

Compound name	Chemical structure	Reference
Vinyl acetate	=	[12]
Methyl acry- late		[26]
Methacryl- amide		[12]
2-Hydroxy- ethyl metha- crylate	Q OH	[60]
Sodium 2-methyl- -2-propene- -1-sulfonate*)	O'Na <sup>+</sup>	[9, 23–25, 28, 32, 40, 44, 47, 48, 55, 57, 59, 67]
Sodium allyl- sulfonate*)	S Na <sup>+</sup>	[26]
[2-(Metha- cryloyloxy)- ethyl]trimethyl ammonium chloride		[41, 68]
3-(Trime- thoxysilyl)- propyl metha- crylate		[22]
2-(Methacry- loyloxy)ethyl phosphate	OH OCO O'OH	[64, 65]
2-Acrylamido- -2-methyl- -1-propanesul- fonic acid	NH VS PO OH	[20, 33, 37]
β-Cyclodextrin with meth- acrylic ester group	OH NH NH	[69]
β-Cyclodextrin with maleic ester group	OH OH OH OH OH OH OH OH OH	[28]

\*) As chain transfer agents in the polymerization.

\_

\_

	Molar ratio		$M_n$	$M_w$	$[M_w/M_n]$	DP
Sample	SamplePoly(ethylene glycol) methyl ether methacrylate $(m)$ Methacrylic acid $(n)$					
PCE-1	1	1.2	23600	40100	1.7	10.8
PCE-2	1	2.4	24150	42760	1.8	10.6
PCE-3	1	3.6	25470	40900	1.6	10.7
PCE-4	1	4.8	25880	41410	1.6	10.4
PCE-5	1	6.0	26460	44980	1.7	10.2
PCE-6	1	7.2	27240	49030	1.8	10.1

T a ble 4. Molecular characteristics of PCE [14]

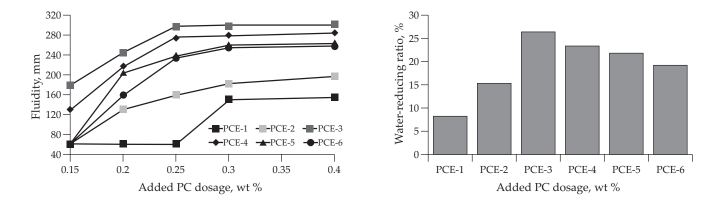


Fig. 3. Effect of carboxylic groups content in PCE on the fluidity at the *w*/*c* ratio of 0.24 (left) and the water-reducing ratios, dosage: 0.2 wt % by cement weight (right) [14]

	PCE 17	PCE 17	PCE 24	PCE 24	PCE 45	PCE 45
	-OCH <sub>3</sub>	-OH	-OCH <sub>3</sub>	-OH	-OCH <sub>3</sub>	-OH
Concentration, wt %	0.16	0.15	0.15	0.19	0.12	0.11

Plank *et al.* [48] modified the poly(ethylene oxide), the side chain of the polycarboxylates, changing its length and kind of terminal group. The authors studied the polycarboxylate superplasticizers with the side chains composed of 17, 24 or 45 repeating units of ethylene oxide, terminated with the hydroxyl (–OH) or methoxy (–OCH<sub>3</sub>) group [Formula (I)]. The effectiveness of the admixtures was evaluated by measuring the content of superplasticizer necessary for achieving a slump (measure of consistence)  $26 \pm 0.5$  cm (Table 5). The admixture with the longest side chains (n = 45), terminated with hydroxyl group appeared to be most effective.

#### PERSPECTIVES OF SUPERPLASTICIZERS DEVELOPMENT

The currently used superplasticizers have some specific advantages as well as weaknesses. There is a growing demand for polymers improving the consistence of the concrete mix with extended time of effective action, and simultaneously inducing no excessive bleeding of water from the mix, air-entraining and set-retardation. Besides the basic criterion of superplasticizer's action, the other requirements are considered, like versatility of using, easy production and application, stable efficiency at various temperatures [71], no negative impact on environment (with a favor for the superplasticizers "consuming" the wastes and allowing for saving the water, cement and energy).

When designing new superplasticizers, the introduction of the acidic group other than carboxylic one into the polymer's structure can be considered. Also, the chain of poly(ethylene oxide) can be modified by introduction of other hydrophilic units, especially those characterized by branched structure, providing reduced viscosity of the polymer solution.

#### REFERENCES

- [1] Czarnecki L., Łukowski P.: *Cement Wapno Beton* **2010**, *15*, 243.
- [2] Aïtcin P.-C.: "Binders for Durable and Sustainable Concrete", Taylor & Francis, London 2008.
- [3] Czarnecki L., Kurdowski W., Mindess S.: "Developments in the Formulation and Reinforcement of Concrete" (Ed. Mindess S.), Woodhead Publishers

Ltd., Abington, Cambridge 2008, pp. 270-284.

- [4] Aïtcin P.-C., Eberhardt A.B.: "Science and Technology of Concrete Admixtures" (Ed. Aïtcin P., Flatt R.J.), Woodhead Publishing, 2015.
- Plank J., Sakai E., Miao C. et al.: Cement and Concrete Research 2015, 78, 81. http://dx.doi.org/10.1016/j.cemconres.2015.05.016
- [6] BS-EN 934-2 (2009), British Standards Institution, London.
- [7] Gołaszewski J.: Construction and Building Materials 2012, 35, 586. http://dx.doi.org/10.1016/j.conbuildmat.2012.04.070
- [8] Plank J., Sachsenhauser B., De Reese J.: Cement and Concrete Research 2010, 40, 699. http://dx.doi.org/10.1016/j.cemconres.2009.12.002
- [9] Plank J., Zhimin D., Keller H. et al.: Cement and Concrete Research 2010, 40, 45. http://dx.doi.org/10.1016/j.cemconres.2009.08.013
- [10] Malferrari D., Fermani S., Galletti P. *et al.: Langmuir* 2013, 29, 1938. http://dx.doi.org/10.1021/la304618f
- [11] Borget P., Galmiche L., Le Meins J., Lafuma F.: Colloids and Surfaces A: Physicochemical and Engineering Aspects 2005, 260, 173. http://dx.doi.org/10.1016/j.colsurfa.2005.03.008
- [12] Duan G., Huang G., Li A. et al.: Journal of Molecular Liquids 2012, 174, 129. http://dx.doi.org/10.1016/j.molliq.2012.07.013
- [13] Winnefeld F., Becker S., Pakusch J., Götz T.: Cement and Concrete Composites 2007, 29, 251. http://dx.doi.org/10.1016/j.cemconcomp.2006.12.006
- [14] Peng X.Y., Yi C.H., Deng Y.H., Qiu X.Q.: International Journal of Polymeric Materials and Polymeric Biomaterials 2011, 60, 923. http://dx.doi.org/10.1080/00914037.2010.551369
- [15] Kjeldsen A.M., Flatt R.J., Bergström L.: Cement and Concrete Research 2006, 36, 1231. http://dx.doi.org/10.1016/j.cemconres.2006.03.019
- [16] Ran Q., Somasundaran P., Miao C. et al.: Journal of Colloid Interface Science 2009, 336, 624. http://dx.doi.org/10.1016/j.jcis.2009.04.057
- [17] Zingg A., Winnefeld F., Holzer L. et al.: Journal of Colloid Interface Science 2008, 323, 301. http://dx.doi.org/10.1016/j.jcis.2008.04.052
- [18] Zingg A., Winnefeld F., Holzer L. et al.: Cement and Concrete Composites 2009, 31, 153. http://dx.doi.org/10.1016/j.cemconcomp.2009.01.005
- [19] Rai S., Gajbhiye N.: Indian Journal of Chemical Technology 2008, 15, 349.
- [20] Üzer E., Plank J.: Cement and Concrete Research 2016, 82, 100. http://dx.doi.org/10.1016/j.cemconres.2015.12.009
- [21] Felekoğlu B., Sarıkahya H.: Construction and Building Materials 2008, 22, 1972.

http://dx.doi.org/10.1016/j.conbuildmat.2007.07.005

[22] Fan W., Stoffelbach F., Rieger J. et al.: Cement and Concrete Research 2012, 42, 166.

http://dx.doi.org/10.1016/j.cemconres.2011.09.006

- [23] Li C., Feng N., Chen R.: *Cement and Concrete Research* 2005, 35, 867.
- http://dx.doi.org/10.1016/j.cemconres.2004.04.031
  [24] Chang W., Li H., Wei M. *et al.*: *Materials Research Innovations* 2009, *13*, 7.
  http://dx.doi.org/10.1179/143307509X402101
- [25] Ma X.: Journal of Wuhan University of Technology-Materials Science Edition 2010, 25, 799. http://dx.doi.org/10.1007/s11595-010-0095-y
- [26] Lu S., Liu G., Ma Y., Li F.: Journal of Applied Polymer Science 2010, 117, 273. http://dx.doi.org/10.1002/app.31984
- [27] Lv S., Duan J., Gao R. et al.: Polymers for Advanced Technologies 2012, 23, 1596. http://dx.doi.org/10.1002/pat.3034
- [28] Lv S., Gao R., Duan J. et al.: Journal of Applied Polymer Science 2012, 125, 396. http://dx.doi.org/10.1002/app.35606
- [29] Lei J., Li H., Du X., Zhang A.: Iranian Polymer Journal 2013, 22, 117.

http://dx.doi.org/10.1007/s13726-012-0110-6

- [30] Lv S., Ju H., Qiu C. et al.: Journal of Applied Polymer Science 2013, 128, 3925. http://dx.doi.org/10.1002/app.38608
- [31] Liu X., Wang Z., Zheng Y. et al.: Materials 2014, 7, 6169. http://dx.doi.org/10.3390/ma7096169
- [32] Li Y., Yang C., Zhang Y. et al.: Construction and Building Materials 2014, 64, 324. http://dx.doi.org/10.1016/j.conbuildmat.2014.04.050
- [33] Zhang Y., Kong X.: Cement and Concrete Research 2015, 69, 1. http://dx.doi.org/10.1016/j.cemconres.2014.11.009
- [34] Łaźniewska-Piekarczyk B.: Construction and Building Materials 2014, 53, 488. http://dx.doi.org/10.1016/j.conbuildmat.2013.11.092
- [35] Lange A., Plank J.: Cement and Concrete Research 2012, 42, 484. http://dx.doi.org/10.1016/j.cemconres.2011.11.017
- [36] Morin V., Tenoudji F.C., Feylessoufi A., Richard P.: Cement and Concrete Research 2001, 31, 63. http://dx.doi.org/10.1016/S0008-8846(00)00428-2
- [37] Chen B.: Journal of Wuhan University of Technology-Materials Science Edition 2015, 30, 533. http://dx.doi.org/10.1007/s11595-015-1185-7
- [38] Ding Q., Zhu Y., Wnag Y. et al.: Journal of Wuhan University of Technology-Materials Science Edition 2012, 27, 768. http://dx.doi.org/10.1007/s11595-012-0545-9
- [39] Ran Q., Liu J., Miao C., Yan S.: Procedia Engineering 2012, 27, 223.
- http://dx.doi.org/10.1016/j.proeng.2011.12.447
  [40] Ng S., Metwalli E., Müller-Buschbaum P., Plank J.: *Cement and Concrete Research* 2013, 54, 191.
  http://dx.doi.org/10.1016/j.cemconres.2013.09.002
- [41] Ren Q., Zou H., Liang M. *et al.*: *RSC Advances* **2014**, *4*, 44018. http://dx.doi.org/10.1039/c4ra05542j
- [42] Jun Y., Jeong Y., Oh J.E. et al.: KSCE Journal of Civil Engineering 2014, 19, 1787.

http://dx.doi.org/10.1007/s12205-014-1189-4

- [43] Liu M., Lei J., Guo L. et al.: Thermochimica Acta 2015, 613, 54. http://dx.doi.org/10.1016/j.tca.2015.05.020
- [44] Kong F., Pan L., Wang C., Xu N.: Construction and Building Materials 2016, 105, 545. http://dx.doi.org/10.1016/j.conbuildmat.2015.12.178
- [45] Ferrari L., Bernard L., Deschner F. *et al.*: *Journal of the American Ceramic Society* **2012**, *95*, 2189. http://dx.doi.org/10.1111/j.1551-2916.2012.05189.x
- [46] Huang H., Qian C., Zhao F. et al.: Construction and Building Materials 2016, 110, 293. http://dx.doi.org/10.1016/j.conbuildmat.2016.02.041
- [47] Wu H., Guo H., Lei J. et al.: Frontiers of Chemistry in China 2007, 2, 322. http://dx.doi.org/10.1007/s11458-007-0060-2
- [48] Plank J., Pöllmann K., Zouaoui N. et al.: Cement and Concrete Research 2008, 38, 1210. http://dx.doi.org/10.1016/j.cemconres.2008.01.007
- [49] Jiang L., Li R., Sun K.: Bulletin of the American Ceramic Society 2013, 5, 852.
- [50] Liu X., Wang Z., Zhu J. et al.: Colloids and Surfaces A: Physicochemical and Engineering Aspects 2014, 448, 119. http://dx.doi.org/10.1016/j.colsurfa.2014.02.022
- [51] Lei L., Plank J.: Cement and Concrete Research 2012, 42, 1299. http://dx.doi.org/10.1016/j.cemconres.2012.07.001
- [52] Lei L., Plank J.: Industrial and Engineering Chemistry Research 2014, 53, 1048. http://dx.doi.org/10.1021/ie4035913
- [53] Sowoidnich T., Rachowski T., Rößler C. et al.: Cement and Concrete Research 2015, 73, 42. http://dx.doi.org/10.1016/j.cemconres.2015.01.016
- [54] Kashani A., Provis J.L., Xu J. et al.: Journal of Materials Science 2014, 49, 2761. http://dx.doi.org/10.1007/s10853-013-7979-0
- [55] Lv S., Gao R., Cao Q. et al.: Cement and Concrete Research **2012**, 42, 1356.
- http://dx.doi.org/10.1016/j.cemconres.2012.06.006
  [56] Ran Q., Wang X., Shu X. et al.: Journal of Dispersion Science and Technology 2016, 37, 431.
- http://dx.doi.org/10.1080/01932691.2015.1042585
- [57] Plank J., Schroefl C., Gruber M. et al.: Journal of Advanced Concrete Technology 2009, 7, 5. http://dx.doi.org/10.3151/jact.7.5

- [58] Pourchet S., Liautaud S., Rinaldi D., Pochard I.: Cement and Concrete Research 2012, 42, 431. http://dx.doi.org/10.1016/j.cemconres.2011.11.011
- [59] Schröfl C., Gruber M., Plank J.: *Cement and Concrete Research* 2012, 42, 1401. http://dx.doi.org/10.1016/j.cemconres.2012.08.013
- [60] Wang D., Cheng P., Xiong W. et al.: Journal of Wuhan University of Technology-Materials Science Edition 2012, 27, 684. http://dx.doi.org/10.1007/s11595-012-0529-9
- [61] Peng X., Li X., Li Z.: Journal of Dispersion Science and Technology 2013, 34, 1265.
- http://dx.doi.org/10.1080/01932691.2012.738130
  [62] Marchon D., Sulser U., Eberhardt A., Flatt R.J.: *Soft Matter* 2013, *9*, 10719.
  http://dx.doi.org/10.1039/c3sm51030a
- [63] Autier C., Azéma N., Boustingorry P.: Colloids and Surfaces A: Physicochemical and Engineering Aspects 2014, 450, 36. http://dx.doi.org/10.1016/j.colsurfa.2014.02.050
- [64] Dalas F., Nonat A., Pourchet S. et al.: Cement and Concrete Research 2015, 67, 21. http://dx.doi.org/10.1016/j.cemconres.2014.07.024
- [65] Dalas F., Pourchet S., Nonat A. et al.: Cement and Concrete Research 2015, 71, 115. http://dx.doi.org/10.1016/j.cemconres.2015.02.001
- [66] Bessaies-Bey H., Baumann R., Schmitz M. et al.: Cement and Concrete Research 2016, 80, 1. http://dx.doi.org/10.1016/j.cemconres.2015.10.010
- [67] Zhu J., Li Y., Zhang G. et al.: Colloid and Polymer Science 2015, 293, 1093. http://dx.doi.org/10.1007/s00396-014-3488-1
- [68] Guo W., Sun N., Qin J. et al.: Journal of Applied Polymer Science 2012, 125, 283. http://dx.doi.org/10.1002/app.35565
- [69] Li Y., Zheng J., Guo H., Lu M.: Journal of Dispersion Science and Technology 2014, 36, 369. http://dx.doi.org/10.1080/01932691.2014.911105
- [70] Wiliński D.: Challenges of Modern Technology 2015, 6, 51.
- [71] Łukowski P.: Journal of Building Chemistry 2016, 1, 31. http://dx.doi.org/10.17461/j.buildchem.2016.105 Received 30 VII 2015. Revised form 10 III 2016.