THE INFLUENCE OF CRUDE OIL PRODUCTS ON RC STRUCTURE DESTRUCTION

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Abstract. It is known that RC industrial structures are liable to the effects of crude oil products, which may have a disadvantageous influence on their exploitation and safety. Within this research, the influence of crude oil products on the physico-mechanical properties of mature concrete, as well as other possible mechanisms of RC progressive failure have been analysed. Crude oil products with a very low neutralisation number were found to be physico-chemical active agents affecting concrete. Experimental analysis undertaken in order to assess the compressive strength of concrete and its bond to reinforcements is described. Long term investigations were conducted in order to determine the effects of a group of crude oil products – mineral oils – on the properties of concrete and reinforced concrete samples. Detailed results of this research are included in the paper. Comparing the influence of different mineral oils on the compressive strength of concrete, leads to the conclusion that there are large differences in their effects. The paper also attempts to answer the question of how the characteristics of RC structure bonds change over time of exposure to mineral oils. The problem of the influence of organic agents on concrete and its bond under conditions of natural humidity is mainly linked to its structure. In the case of oil products, there are four possible mechanisms i.e., biological, chemical, physical and physico-chemical, affecting concrete, its bond and, sometimes, reinforcements. Each of these mechanisms is presented. Necessary laboratory tests were performed in order to prove the thesis that the content of active surface polar molecules in active agents is the reason behind concrete damage brought about by crude oil products. The conducted experiments demonstrated that the effects of artificial environments (on concrete) prepared by the author, are the same as those of widely used industrial mineral oils.

Keywords: RC structures, influence of crude oil products, concrete destruction mechanisms, compressive strength, bond stress, surface active polar molecules.

1. Introduction

In technical literature, the effects of crude oil products on concrete are classified either as non-harmful or only mildly harmful; however, there is evidence that serious damage can occur as a result of such products (Миловидов, Алексеева 1970; Pluta et al. 1980; Khoury 1981; Błaszczyński 1995; Onabolu 1986). Oil products affect tank walls, foundations and other constructions supporting machines, and storage floors, therefore, the range of the problem is wide and the subject should be considered of great technical importance, especially when regarding industrial construction. Inspection of frame foundations used for large power turbines (Fig. 1) showed that, with the passing of time, considerable damage can be observed. The damage, mainly cracks and fissures, and a decrease of the structure’s dynamic stiffness, often causes serious damage to the turbines. It was also noted, that strong oiling of the foundations is one of the main factors causing these damages (Błaszczyński et al. 1985).

Near the bottom slab of the foundation, a water-oil environment, which causes damage through the biological mechanism of destruction, can be observed (Fig. 2).

Fig. 1. Observed damage of a turbine’s frame foundation

The oiling of structural elements is a problem also present in industrial buildings. Strongly oiled, reinforced, concrete floors display considerable damage, which often impairs a building’s exploitation. After a few years, significant enlargement of cracks brought about by the influence of oil can be observed (Миловидов, Алексеева 1970; Pluta et al. 1980). Biczok (1972) presented an example of an oiled, reinforced, concrete floor in which
significant cracking and deflection occurred and, during repair work, concrete was easily separated from the reinforcement bars (Fig. 3a). The same problem was noticed in the floors of industrial buildings (observed at the Social Psychology University in Warsaw, recently analysed by the author) (Fig. 3b).

Biczok (1972) noted that concrete specimens taken from the oiled floor demonstrated a decrease in strength of up to 40%. This was also shown in Manns and Hartmann’s as well as Wasiliew’s research (Manns and Hartmann 1977; Васильев 1981). On the other hand, Grabiec (1973) presented contradictory results regarding oiled concrete; in his research, the relatively long-term oiling of samples (about 5.5 years) did not cause a decrease in compression and tensile strength below the initial values. During his investigations, Grabiec used various mineral oil types: Lux 10, Wrzecion 2 and TZ-4 water-oil emulsion (5:1). Similar results were obtained by Dominas (1970) for Wrzecion 2 oil. Nevertheless, the latest research by Runkiewicz et al. (2002) demonstrates that oiling concrete with mineral oils adversely affects the mechanical characteristics of concrete. The researchers used machine oil (L-AN 22) and cooling emulsion (EMULGOL ES12), with the period of research lasting 1260 days (almost 4 years).

The influence of Hydrocarbon-products (crude oil products) over a long period of time in comparison to water was presented by the author in earlier research (Blaszczyński 1994). The results of these studies varied greatly: from a lack of influence observed in the case of kerosene, petrol, or vaseline oil, to significant changes caused by certain mineral oils.

The author’s longstanding research focusing on the influence of mineral oils on mortars and cement-based concretes, has led to rebuttal of the thesis concerning the considerable influence of the viscosity of crude oil products, especially mineral oils, on the degree of concrete destruction, and enabled one to discriminate between more and less destructive substances from the mineral oils group; such findings contradict a common thesis that assumes equal affects of all mineral oils (Blaszczyński 1994, 2002).

Current research pertaining to the influence of mineral oils on the adhesion of plain reinforcements to oiled concrete was conducted by Runkiewicz et al. (2002). His investigations, conducted by means of the pull-out method, incorporated plain bars (steel St0) of Ø 14 mm. After about 4 years of subjecting the given material to the effects of mineral oils, a fall of maximum adhesion from 17% to 33% was reported.

2. Methods, materials and results

2.1. Influence of mineral oils on concrete

Long-term laboratory experiments have been conducted to assess changes in the physico-mechanical characteristics of concrete contaminated by oil. The compressive strength was determined, with the use of 100 mm cubes and in accordance with PN–EN 12390–1:2001, for concrete type – B25, which is the most commonly used concrete type for industrial RC structures in Poland (5 specimens, standard deviation $s_{fc} = 0.84 – 2.87$, variation index $\nu_{fc} = 2.25\%$ – $6.99\%$). The average 28 day compressive strength of concrete equalled $f_{cm} = 29.8$ MPa. The water-cement ratio was 0.59, while the aggregate-cement ratio equalled 6.70. Three different oils were used in the research: TU–20, M–40 and H–70 with respective viscosities of $81.0 \text{ mm}^2/\text{s}$, $211.4 \text{ mm}^2/\text{s}$, and $383.0 \text{ mm}^2/\text{s}$, at $20 ^\circ\text{C}$. These oils have low neutralisation numbers with values between 0.05 and 0.075 mgKOH/g. The first application of oils to concrete took place 2 months after casting; subsequently, the specimens were examined every 4 months (up to the 36th month), and every 12 months (from the 36th month onwards) over a period totalling 72 months. The control specimens (samples) were additionally examined 28 days, and 2 months after the application of oils. The concrete was oiled two months after casting, which marks the beginning of the...
time of oiling \( t_z \), when the average compressive strength of concrete \( f_{cm} = 37.35 \text{ MPa} \). Comparison of the influence of oils on compressive strength shows that oils TU–20 and H–70 decrease \( f_{c}^{b} \) the most. Changes in the compression strength of concrete \( f_{c}^{b} \) are presented in Fig. 4.

The results illustrate, that the significant decrease of \( f_{c}^{b} \) occurs as a result of contamination with oils. Comparison of the influence of oils shows that TU–20 and H–70 oils are responsible for the largest decrease in \( f_{c}^{b} \). When comparing the effects of oils used it should be noted that the final \( f_{c}^{b} \) values of concretes exposed to oils differ significantly from the control group, and range from 55% (H–70 oil) to 10% (M–40 oil). While a considerable decrease in \( f_{c}^{b} \) occurred after the application of H–70 and TU–20 oils, M–40 oil (of medium viscosity) did not cause a decrease in \( f_{c}^{b} \) below the initial value.

2.2. The influence of mineral oils on concrete bond

Conventional reinforced concrete structures are designed under the assumption that concrete and steel bars work together in carrying loads. The collaboration of steel and concrete, based mainly on adhesion (natural adhesion) and friction (in the case of plain bars), as well as mechanical bond (in ribbed bars), is a distinctive feature of reinforced concrete. In the case of plain bars, with a surface roughness coefficient of \( Ra \leq 0.1 \mu \text{m} \), the value of the bond depends only on natural adhesion. A solution to the problem of the influence of mineral oils on the natural adhesion of concrete to plain reinforcements was sought through experimental research conducted on specimens of the author’s own design (Fig. 5).

The character of changes in the case of the influence of crude oil products on the bond between ribbed bars of different ribbed ratios \( f_{R} \) (Manns and Hartmann 1977) and concrete, differs.

Changes in the natural adhesion of investigated concretes to plain bars over time following oiling with TU–20 mineral oil is illustrated in Fig. 6.
Rangan 1998), and the influence of different products (Blaszczynski 1995). However, the behavior of such specimens differs greatly from that of a cracked concrete structure (Pędziwiatr 2008).

Changes over time in the bond stress of analyzed concretes and ribbed bars, following oiling with TU–20 mineral oil and tested for by means of the pull-out method, are shown in Fig. 7. The decrease in bond is gradual and, as in the case of concrete compressive strength, takes six years to stabilize (Fig. 4).

2.3. Investigations incorporating artificial, physico-chemically active environments

In order to prove the physico-chemical mechanism, an artificial environment was prepared. It was composed of a mix of vaseline oil (basic oil – not active) and oleic acid (improver – surface active fluid). 100 mm concrete cubes, made of grade C20/25 concrete were exposed to four different artificial environments i.e., vaseline oil, vaseline oil with 1% oleic acid, vaseline oil with 5% oleic acid and vaseline oil with 10% oleic acid, six years after casting. Subsequently, the compressive strength of concrete was examined over a period of 4 years, as presented in Fig. 8.

As illustrated in Fig. 8, only vaseline oil is non-harmful to concrete. The addition of oleic acid (surface active molecules) has a similar effect on the compressive strength of concrete as the analyzed mineral oils (Fig. 4). The intensity of the penetration of two artificial environments i.e., vaseline oil and vaseline oil containing 10% oleic acid, was also examined by measurement of the increase in the values of mass (Fig. 9). Nearly complete saturation of samples occurred within a period of 12 months. After this time, the decline in strength could be caused only by the destructive processes of the added surface-active substance (oleic acid). Significant differences were not observed between the kinetics of saturation and the increase in mass in either of the applied environments. Thereby, it was confirmed that physical phenomena, such as the heave of penetrating material or its filling in of gaps, were not the reasons behind changes illustrated in Fig. 8.

3. Concrete and its bond destruction mechanisms caused by crude oil products

The reasons behind the destruction of concrete structures are usually complex, nevertheless, a dominating cause can generally be established. In the case of crude oil
products, there are four possible mechanisms of concrete destruction i.e., biological, chemical, physical and physico-chemical. The same mechanisms damage bonds between concrete and steel reinforcements, and are due mainly to the properties of concrete.

3.1. Biological and chemical mechanisms

In water-oil systems, two groups of bacteria develop: aerobic oil oxidizing bacteria – OOB, and anaerobic sulphate reducing bacteria – SRB (Morgan et al. 1983). The first group grows on crude oil molecules, while the later, on water molecules. OOB growth on crude oil molecules appears earlier than SRB development in water containing crude oil molecules, which suggests that a form of nourishment symbiosis is present between them (Dahan 1984).

OOB produce organic acids, mainly acetic acid (CH₃COOH) and, in effect, lower the pH of water. SRB produce hydrogen sulphide (H₂S) and do not cause a decrease in pH values (Fig. 10 and Postgate 1979). The extent of their growth and the level of H₂S production are directly connected with the presence of crude oil and its products (Wilkinson 1983).

In the case of crude oil, the presence of 50–100 mg/l H₂S was reported (which is 3–5 times higher than the limiting standard value), with pH values decreasing from 8.4 to 6. Sulphur bacteria i.e., Thiobacilli X and Thiobacilli Concretivori, present under these conditions, cause the oxidation of hydrogen sulphide, production of sulphuric acid (H₂SO₄), and a decrease in pH values of 2, 3, and even 1.

Onabolu et al. (1985) state that this acid is not intercepted from water by crude oil, even at a very low pH value (2.2) and high temperature of the environment (80 °C).

The following factors are responsible for the induction of chemical destruction of concrete structures, in an environment of crude oil products:

- a decrease in pH value of the oil-water environment, caused by the presence of aerobic bacteria and sulphate bacteria, existing in an anaerobic bacteria environment;
- organic acids produced by aerobic bacteria (acetic acid), and acids remaining after the refining process of crude oil, or created as a result of the exploitation of its products (machine and engine oils);
- a hydrogen sulphide water solution produced by anaerobic bacteria;
- sulphuric acid produced by sulphate bacteria;
- sulphur and its compounds contained in crude oil and its products.

In summary, two kinds of chemical destruction mechanisms are dealt with: a generally acidic (pH value decrease of the oil-water environment, organic acids) and sulphate (sulphur and its compounds).

The effects of acids on concrete include reduction of its strength, a decrease in weight, an increase in cement stone dissolution or porosity, and the exposure of aggregate grain (Czarnecki et al. 1994). The degradation process depends on the pH value of the environment. Destruction is to be expected at pH values below 6.5; for instance, according to Romben (1978), concrete destruction at pH values between 3 and 6 proceeds in an extent proportional to the square root of action time.

The effect of organic acids on concrete, similar to that of non-organic acids, is based on the reaction with calcium hydroxide and, partly, with aluminium compounds. Acetic acid, produced by bacteria, reacts with free hydroxide or calcium oxide, creating soluble calcium acetate (Kawai et al. 1997) which, in turn, causes calcium loss; therefore, even thinned solutions are strongly aggressive. Other carboxyl acids (for example oleic acid or stearic acid) react with Ca(OH)₂, creating soft, insoluble lime soaps (calcium oleate or calcium stearate). If there is a large inflow of these acids, after Ca(OH)₂ runs out in the cement, decomposition of calcium silicates can take place, along with silicate gel precipitation and the formation of further calcium soaps. This leads to the softening or cracking of concrete (Gruener 1983; Ściślewski 1999).

The acid content of crude oil and its products is specified in its neutralization number, which equals the amount of KOH (in milligrams) needed to neutralize unbound acids present in 1g of the given product (PN-85/C-04066). According to Steinbach (1967) and Weiss (1961), chemical reactions of non-vegetal oils can be expected only if the neutralization number is greater than 0.25 mg KOH/g. This value is alleged to be the chemical limit of concrete structure destruction caused by crude oil products (Vorläufiges Merkblatt... 1966). However, in research involving oiled concrete (Manns and Hartmann 1977) it was reported, that despite the use of exploited machine oil (neutralization number equal to 0.48 mg KOH/g), no chemically induced damages were observed. The damages were of physico-chemical origin. In more recent works, the limiting value has been increased to 0.5 mg KOH/g (Tscherner 1986), or even 1.0 mg KOH/g (Brauer 1985). Nevertheless, in practice, it is recommended to employ the safest (initial) value of 0.25 mg KOH/g. Below this value, the presence of polar organic
acid molecules (including the – COOH group) in crude oil products affect the concrete structure physico-chemically more so than chemically. This does not mean, however, that organic acids with a neutralization number $<0.25$ mg KOH/g do not cause the chemical reactions described above, but rather that the quantity of products (lime soaps) resulting from these reactions does not lead to corrosion and structural changes in the concrete.

The chemical influence of mineral oils was investigated by Wasiliew (Васильев 1981). He proved that organic acids, present in these oils, react chemically with Ca(OH)$_2$ and, thus, create soft lime soaps, causing the softening of concrete, and destruction of samples after just one and a half years. The research did not specify what neutralization number characterized the examined specimens.

For experimental purposes, the influence of various acids on the acidity rate of crude oil was tested. Fig. 11 illustrates the effects of organic (acetic) and non-organic (nitric) acids, as well as two organic (acetic and oleic) acids on oil acidity. When the content of acetic acid reaches 0.05%, crude oil becomes an aggressive environment for concrete, while at 0.5% it has a neutralization number of 5.12 mg KOH/g and, therefore, creates a very aggressive environment. The addition of non-organic acid did not cause crude oil acidity to change (Fig. 11a) (Khoury et al. 1985).

Comparison of high- and low-carbon, organic acids allows one to conclude that their acidic characteristics decrease along with an increase in the amount of carbon atoms in the compound (Fig. 11b). From the author’s research on the influence of the addition oleic acid on crude oil and the neutralization number of its product, it can be stated that when the content of this acid significantly exceeds 10% of the volume, the lowest safety limit (0.25 mg KOH/g) for the chemical destruction mechanism is surpassed. In practice, however, the content of this type of acid does not exceed a few percent.

It can, therefore, be stated that concrete corrosion caused by organic acids can occur only under the influence of acetic acid and is not to be expected in the case of crude oil products containing high-carbonic acids (oleic and stearic acids).

Acetic and sulphuric acids are classified as factors causing fast corrosion of concrete. Over extended periods of time (a few years), even H$_2$S causes detrimental changes (Gruener 1983). Chemical reactions between concrete and sulphuric acid, as well as other sulphur compounds, result in the formation of compounds containing soluble calcium hydrogen which, in turn, makes concrete weaker and more porous (Piasta, J. and Piasta, W. G. 1994).

When acids or their salts solutions are able to reach the reinforcement through pores and cracks, corrosion of reinforcements occurs, causing further cracking and peeling of concrete.

### 3.2. Physical and physico-chemical mechanisms

In the case of liquids, the physical destruction mechanism generally occurs as a result of internal pressure caused by bulking which, in turn, usually takes place due to changes in temperature. However, during the exploitation of crude oil products, temperature changes range from 15 °C to 80 °C, which does not result in pores and micro-cracks expanding notably and, thus, is not followed by an increase in internal pressure.

Extensive studies of literature, as well as the author’s own research, did not confirm any signs of concrete damage by crude oil products, stemming from the physical mechanism of destruction.

The phenomenon, which can be termed “the physico-chemical mechanism of concrete destruction by organic substances”, is compromised of three basic elements:

- a) surface energy reduction in the concrete’s structure due to active substance absorption;
- b) the wedging effect;
- c) the insufficiency of concrete strength due to its hydrophobization, caused by the lack of hydration.

![Fig. 11. The influence of organic and non-organic acid content on oil acidity: a) acetic acid and nitric acid (Khoury et al. 1985); b) acetic acid and oleic acid](image-url)
Relevant substance absorption can occur only when connected with a decrease of free energy in this layer, hence, along with a drop in interfacial tension. The ratio of surface tension to concentration in a balanced state was labelled as absorbing substance surface activity by Re-binder (Пебиндер 1966), which characterizes the absorbing substance’s ability to change the interfacial tension of the absorbant. Substances which decrease interfacial tension are called surface-active substances.

A substance can be considered surface-active if its molecules are asymmetric and contain polar hydrophilic groups and non-polar hydrophobic groups. The molecules of surface-active substances become more absorbent, the more distinctively marked the asymmetry of hydrophobic groups and the larger the differences in polarity of both phases (the absorbing substance and the absorbant) comprising the adjacent surface.

Forces aimed at decreasing free surface energy by decreasing surface tension, act on the adjacent surface. The decreasing surface energy causes the structure to loosen, enabling physico-chemically active substances to penetrate it. Due to the surface and internal stability of the structure of crystalline substances, the only means of decreasing the surface energy and saturation of forces acting the surface of such substances, is to tie extraneous molecules from the environment. The decrease in the surface energy of a solid caused by the saturation of its internal space with molecules of a polar liquid is usually much lower than the surface energy of the liquid itself (Robertson and Mills 1985).

In the case of non-polar liquids, such as benzene or paraffin, a loss of strength is not observed, since, due to their non-polarity and the diameter of their molecules (which is 2–3 times larger than that of water molecules), they are unable to effectively penetrate the porous cement structure. Smaller crude oil fractions, in which polar molecules are smaller than water molecules, lose their ability to penetrate gel pores by means of molecular sieve.

The mechanism of adhesiveness between the cement matrix and aggregate, as well as the cement matrix and reinforcements, and the coinciding operating contact zone destruction caused by crude oil products, are most likely connected with the wedging effect, which occurs on the liquid-solid interface. This phenomenon has been known for years in metal technology, and used in their processing processes. Re-binder and Lichtman obtained interesting data during their research focusing on the influence of surface active molecules contained in lubricants on metal processing procedures, especially sawing and turnery. In their works (Лихтман et al. 1954, 1962), they proved that noticeable softening of metal surfaces can be observed in the presence of polar molecules. In this environment, the increased softening effect has an energy-forced character and is caused by the surface energy of a solid. When a micro-crack is present on a solid’s surface, polar molecules penetrate it, along the walls towards its tip, until they reach the point at which the crack’s dimensions prevent further penetration (Fig. 12).

As the collision between the molecules and the surface increases, wedging pressure (marked in the figure by arrows) occurs, resulting in the weakening of bonds and widening of cracks. Due to interaction of the solid molecules and surface-active molecules, further migration of the later can lead to the slow (without internal energy loss) splitting of solid molecules. The internal splitting kinetics of a structure is determined by the penetration speed, from the surface-active substance to the deteriorated surface, of polar molecules. The dispersed molecules, moving into the newly created (as an effect of wedging) spaces of micro-cracks, prevent the cracks from closing, which accelerates the destruction process. The internal cracking progress is defined by the movement speed of molecules along newly created surfaces towards the new apex of the crack. The splitting action of solutions containing surface-active molecules is considerably stronger than that of basic substances, and has no direct connection with its viscosity.

The wedging effect has been proven by microstructural research, conducted by Onabolu (1986), on concrete samples saturated with crude oil. In Fig. 13a destruction, including progressive cracking and adhesiveness between the aggregate and cement stone, can be observed. Results of the wedging effect are also visible in Fig. 13b, which presents a piece of loose concrete from a reinforced concrete element.

Each splitting causes the surface-active molecules to deposit on the walls, becoming eliminated from the penetrating substance. At some point, a critical saturation of surface-active molecules is reached (all surface-active molecules are eliminated from the oil phase), which stops the further decrease in strength. Therefore, in the event of too few surface-active molecules water that isn’t surrounded by them is able to act upon and bond to the cement stone structure. If an organic polar substance is not delivered simultaneously, water molecules begin to prevail. A similar migration process follows but, this time, with a water system which causes renovation and strengthening of the bonds, and leading to an increase in concrete strength. This phenomenon explains the various effects observed in studies regarding the influence of mineral oils.

The influence of M-40 oil, presented in Fig. 3, is caused by the small amount of surface-active molecules present in its composition, as a result of imperfections in
the refining process of its oil base. Machine oils are one-
fraction hydrocarbons, containing a very small amount of
organic surface-active improvement additives. Results for
Wrzecion 2 oil, similar to those for M–40, obtained by
Grabiec (1973) and Dominas (1970), can be explained by
the fact that it is the same machine oil, but with a consid-
erably reduced viscosity (owing to its use in high-speed
spindles). The effects of machine oils have been estab-
lished by the latest research conducted by Runkiewicz et
al. (2002), during which L–AN 22 machine oil was
tested. The above research stands as evidence that oil
viscosity does not lead to changes in the physico-
mechanical features of concrete and reinforced concrete.
Although the values of kinematical viscosity of the ana-
alyzed oils differed vastly at a temperature of 50 °C
(Wrzecion 2–8 mm²/s, L–AN 22 – 22 mm²/s, M–40 –
40 mm²/s), their influence was comparable.

Turbine (TU–20) and hydraulic (H–70) oils are
multi-fractional compositions, of varying refining rates
and distinctive amounts of improvement additives. Hy-
draulic oils are composed of oil bases with a medium
refining rate, containing a small amount of polar con-
taminants, while turbine oil bases are well purified.

Lux 10 oil, used by Grabiec (1973), is non-
improvement oil, containing small amounts of polar con-
taminants. It contains polar remains – creosols and fur-
furyal alcohol, acquired during the production process.
The transmission of Lux 10 oil through a column contain-
ing silicone gel caused the separating of these substances
along with the absorption of a portion of the aromatic
hydrocarbons into the column’s contents. The quantity of
these substances did not exceed 0.01%, which explains
such small decreases in strength which later tended to
increase, observed in Grabiec’s experiments.

As a whole, the phenomenon of the physico-
chemical influence of organic substances depends on the
competition between organic surface-active molecules
contained in these substances with a liquid the polarity of
which is comparable to that of water. The competition, in
proportion to the positive influence of water, can be cal-
led the internal hydrophobization of concrete structures,
because molecules containing long hydrocarbon-chains
hydrophobize the cement stone, preventing water molecu-
les from penetrating the created limiting layer, thus, im-
peding cement grain hydration. This was proven by deri-
vatographic research conducted on concrete samples
oiled with TU–20 oil (Kozarczewski 1987). In the above
studies, it was stated that the presence of C–S–H phase
contents in TU–20 oil is smaller than under normal matu-
ring conditions. The comparison of derivatograms (peak
height) of non-oiled and oiled samples reveals that in the
case of the later ones the C–S–H phase is restricted; these
findings were also confirmed by Onabolu’s microstructu-
ral research (Onabolu 1986).

Fig. 14b presents the microsection of a sample which
had been soaked with crude oil. The lack of certain crystal-
line phases is clearly visible in comparison with the micro-
section of a sample not treated with crude oil (Fig. 14a).
Due to the hydrophobization process, internal drying of concrete can occur, restricting the strengthening process and, thus, resulting in the occurrence of contractile micro-cracks which can be penetrated by surfactant molecules.

The comparison of the physico-chemical influence of water to that of crude oil derivatives allows one to draw conclusions regarding the harmfulness of surfactant (polar) substances containing hydrocarbon-chains. The detrimental effect of the hydrocarbon-chain occurs after it is joined to the hydrophilic polar part. This gives an explanation as to why kerosene, petrol, crude oil, diesel oils and other related substances, which consist of short-chained hydrophobic parts, do have such a strong impact. In the case of fuels, water is the main polar substance, while other, also short-chained, polar molecules occur in small amounts.

In concrete, the main place of physico-chemical interference is the area of adhesiveness between the cement matrix and the aggregate and, in case of reinforced concrete, additionally the area between reinforcements and cooperating concrete. The destruction extends from the largest areas (large grain aggregate adhesive area) to the smallest ones (small grain aggregate – sand – adhesive area). Faiyadh (1980) showed that stress loss caused by the influence of crude oil on the same classes of concrete is larger for concrete containing 20 mm aggregate than of that, which contains 10 mm aggregate. Koroleva (1993) believes that, with the lack of satisfactory aggregate and cement matrix adhesiveness, the compression strength of some concretes can decrease up to 14% and, in some cases, even up to 20%. The quality of the contact zone influences not only concrete strength and deformability, but, to a great extent, its durability as well. This area increases considerably in the case of high-strength concretes. Contamination of the aggregate with various substances, such as oils, asphalts and other organic matter, significantly decreases the energy of the contact area net. A thin layer of paraffin wax, covering the surface of thick granite aggregate, reduces concrete compression strength by 30–35%, and its tensile strength by 37%. In the case of limestone aggregate, these values are twice as large as those of Flaga (1971). The same phenomenon occurs when the aggregate is contaminated with a thin layer of clay. This significantly impacts the size of the contact area, and leads to a decrease in concrete strength, crack resistance and durability. Contamination of an aggregate’s surface can reduce concrete bending strength by 10–20% while lowering its fatigue strength fourfold.

When comparing the influence of water and organic molecules over time, it can be stated that, in case of water, an initial dramatic decrease in strength is followed by its increase, which attests that the positive chemical influence of water on the cement matrix is stronger than its negative, physico-chemical effect. At the same time, the influence of surface-active substances present in mineral oil derivatives has an adverse effect (Fig. 1), mainly due to the wedging effect, which does not occur in the case of water, but can occur in the contact zone between concrete and steel bars.

The negative influence of RC structure oiling with crude oil products, over time, applies not only to concrete structures and the contact zone, but to reinforcements as well. Fortunately, crude oil products are not harmful to reinforcements until they become biologically or chemically active. In the event of biologically or chemically active environments, the protection of reinforcing steel from corrosion, e.g. by sulfur polymer composites, becomes necessary (Hola and Książek 2009).

Blaszczyński and Sciallo (2006) presented a method of evaluating the limiting capacity of a reinforced concrete element exposed to the influence of mineral oils.

4. Conclusions

The influence of crude oil products can have a very negative effect on the compressive strength of concrete, as well as its bond. Comparison of the influence of oils on compressive strength shows that TU–20 and H–70 oils decrease \( f_\text{ck} \) the most. When comparing the effects of the analyzed oils it should be noted that the final \( f_\text{ck} \) values of concretes exposed to oils differ significantly from the control group, and range from 55% (H–70 oil) to 10% (M–40 oil). The largest decrease was observed in the case of H–70 and TU–20 oils, but M–40 oil (with medium viscosity) did not cause \( f_\text{ck} \) to decrease below the initial value. The change in bond stress of investigated concretes to ribbed bars over time after oiling shows that the weakening of bond is gradual and becomes stabilised after 6 years, as in the case of concrete compressive strength (Fig. 4).

On the basis of comparative analysis one can, with large probability, ascertain that artificially created physico-chemical environments cause changes comparable to those induced by universally applied industrial mineral oils. The lack of essential differences in the penetration intensity of vaseline oil and vaseline oil with 10% oleinic acid, as well as on slight changes of the compression strength among the environments created from vaseline oil and vaseline oils containing 10% and 5% oleic acid prove that in the case of industrial mineral oils, the physical mechanism of destruction does not occur.

The physico-chemical destruction process of concrete structures is complex and ambiguous. Besides areas where hydration and carbonation processes take place, there are other areas in which such processes are restricted, as well as areas in which the wedging mechanism occurs. The concrete deterioration mechanisms present model situations (in which each occurs individually). In practice, however, all presented mechanisms can occur simultaneously, in parallel time, but at different intensity, or successively, one following another. Both in parallel and in serial time, these mechanisms influence one another through interactions between their products and by chemically and physico-chemically induced changes in the concrete structure, which affect the proceeding phenomena.

For the first time, all destruction mechanisms caused by crude oil products were introduced and systematised. The
physico-chemical mechanism is most common, while the biological mechanism can occur only in a water-oil environment, which exists in the case of industrial structures.

Protection to crude oil products could be done by surface coating of RC elements. Only not cracking coatings are successful in case of crude oil products influence. Interesting modeling of corrosion protection for reinforced concrete structures with surface coatings is presented in Kamaitis (2008, 2009).

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*PN-85/C-04066. Przetwory naftowe. Oznaczanie liczby kwasowej i zasadowej oraz kwasowości metodą miareczkowania*
NEAPDIRBOTOS NAFTOS PRODUKTŲ ĮTAKA GELŽBETONIO STRUKTUROS SUARDYMUI
T. Z. BLASZCZYŃSKI
Santrauka
Yra žinoma, kad gelžbetoninės pramoninės konstrukcijos yra jautrios neapdirbotos naftos produktams, o tai gali turėti neigiamą įtaką jų eksploatacijai ir saugumui. Šiame tyrimo analizuojama neapdirbotos naftos produktų įtaka brandaus betono fizinėms ir mechaninėms savybėms. Aptariamai galimi poveikiai, skatinantys progresyvų gelžbetonio irimą, taip pat neapdirbotos naftos produktų grupės – mineralinių naftos produktų – poveikį betono savybėms ir armaturos bandiniams. Šiame straipsnyje pateikta informacija apie galimus poveikius, skatinantys progresyvų gelžbetonio irimą, taip pat neapdirbotos naftos produktų grupės – mineralinių naftos produktų – poveikį betono savybėms ir armaturos bandiniams.