Accelerated aging studies of the selected commercial films

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Abstract: The Arrhenius method was used to investigate the effect of accelerated aging conditions on the physical and mechanical properties of polymer films for technical, medical, food and office applications. Temperature and UV radiation were taken into account as environmental factors. It was shown that the tested films are more sensitive to thermal degradation than UV radiation. The reduction in tensile strength after thermal aging does not exceed 40%, and in the case of UV radiation, 14%. The estimated lifetime of the tested films is over 5 years.

Keywords: films, mechanical properties, accelerated aging, Arrhenius method.

Badania przyspieszonego starzenia wybranych folii handlowych

Streszczenie: Metodą Arrheniusa zbadano wpływ warunków przyspieszonego starzenia na właściwości fizyko-mechaniczne folii polimerowych do zastosowań technicznych, medycznych, spożywczych i biurowych. Jako czynniki środowiskowe uwzględniono temperaturę i promieniowanie UV. Wykazano, że badane folie są bardziej wrażliwe na degradację termiczną niż promieniowanie UV. Zmniejszenie wytrzymałości na rozciąganie po starzeniu termicznym nie przekracza 40%, a w przypadku promieniowania UV, 14%. Oszacowany czas życia badanych folii wynosi ponad 5 lat.

Słowa kluczowe: folie, właściwości mechaniczne, przyspieszone starzenie, metoda Arrheniusa.

Plastics production has increased more than 20-fold during the past half-century. In 2015 alone, global plastics production was about 322 million tons, and reached 335 tons in 2016 [1, 2]. Due to the low cost, transparency, durability and easy processing, polymers have been widely used in agriculture, industry, and human life. The disadvantages of plastics are low resistance to high temperature and poorer mechanical properties, compared to, e.g. metals or ceramics.

Films are made of polymers or non-ferrous metals up to 2 mm thickness. Plastic films are most often made by the following methods:

– blow moulding extrusion (molten polymer is extruded through the slot in a circular cross-section, and air pressure makes it form a sleeve),
– cast extrusion (molten polymer is extruded through a flat nozzle and cast on a cooled mould).

Films may be single or multi-layer. In the second case, the film consists of different layers of materials, thanks to which it combines the features of components. Films may also be combined with paper or aluminum. Polymers used for films are most often polyethylene (PE), polypropylene (PP), ethylene copolymers with vinyl acetate or vinyl alcohol, ethylene copolymers with methacrylic acid, polyvinyl alcohol and fluoropolymers. Properties that determine the durability of films are characterized by numerous indicators, physical values. Amongst the required properties are:

– technological (linear dimensions, composition, mass per unit area, manufacturing defects, colour, printed design, gloss),
– physical (composition, mass per unit area, density, change of dimensions – shrinkage, flatness deviation, flexibility),
– tensile [strength and elongation, tear resistance, resistance to puncture (free-falling dart method), adhesion, surface coefficient of friction, weld strength, resistance to separation of layers],
– bio-physical, ecological and hygienic (water absorption, migration of dyes, transmission of odour and flavour, penetration of water steam, water permeability, oxygen penetration, fat penetration, monomer content),
– special (susceptibility to formation, thermal resistance, wettabiliy, electrostatic properties),
– barrier (resistance to microorganisms, UV radiation, high temperature),
– sensorial,
– aesthetic.

Films are used as materials with a long durability period. Depending on the type, the period of storage in appropriate conditions (temperature below 25°C, no access of moisture, light, oxygen, ozone, and chemical...
substances, position preventing deformations) is 3–25 years. However, the precise forecasting of the durability period under conditions of use, the so-called lifetime is very important for safety reasons. Changes can result from both chemical (chemical degradation) and physical factors (physical degradation). The service life is defined as the service life after which the material reaches the threshold level of the test value (usually 50% of the initial value) at the service temperature. For the Arrhenius method, this value is determined by extrapolating a linear relationship to the service temperature. In literature, the other terms can also be found [3, 4]:

– storage life, shelf life, *i.e.* the maximum period from the date of cross-linking to the date of installation (the first use of the product),

– service life, *i.e.* the expected lifetime or the acceptable period of use (manufacturer's warranty).

The storage life depends on conditions in which the product is stored.

Over time, the physical properties of the film can change in different ways and, as a result, become useless due to excessive hardening, softening, breaking or other surface damage. Changes in physical properties may be caused by environmental factors, such as oxygen, ozone, light, and increased temperature.

Artificial accelerated aging tests, which usually expose materials to artificial light sources and simulated environment conditions, are commonly conducted to evaluate the weatherability and service life of polymeric materials [5, 6].

Changes in physical properties also depend on the chemical composition of the film [7, 8].

The aim of the study was to determine the influence of factors related to the accelerated aging process, taking into account temperature or UV radiation as environmental factors, on the maintenance of the physical and mechanical properties of films. Additional effects included the selection and determination of the physical properties of films and aging conditions in the context of determining the temperature, exposure time, UV radiation dose, as well as the final determination of the impact of the accelerated aging simulation process on the change of selected physical and mechanical properties of the films before and after pre-set time intervals.

**EXPERIMENTAL PART**

**Materials**

The plastic films for various purposes were tested, *i.e.*:

– technical film on anti-flooding sleeves (two types): sample 1 – anti-flooding sleeve Z 2 and sample 2 – anti-flooding sleeve W 2


– food film (one type): sample 6


The basic physical properties of the tested films (sample 1–7) are presented in Table 1.

**Methods**

Mass per unit area, thickness and density were determined according to PBM-53/ITB procedure “Plastic Foils research methods”. The test of mass per unit area consisted in determining the mass of the samples with a specific area (100 cm²), and then expressing them in g/m² with accuracy to three significant figures. The thickness of the samples was measured with an accuracy of 0,01 mm at a specific measuring pressure 0,5–1,0 N and a measuring foot with a specific diameter 2,5–10,0 mm. The density was calculated using the mass per unit area of the samples and their thickness according to the formula \( \rho = m/V \), where \( m \) is the mass of the sample and \( V \) – its volume.

Tensile strength was measured with an Instron universal testing machine according to PN-EN ISO 527-3:2019-01 standard.

Exposure to environmental factors such as temperature and UV radiation were determined according to ASTM F1980-16, PN-EN ISO 291:2010 standard and PBM-53/ITB:2019 procedure.

The tests were carried out on conditioned samples after the accelerated aging, taking into account the following time intervals: 3 years and 5 years.

The aging simulation included an incubation at temperature and the effects of ultraviolet radiation. Plastic films acc. to the requirements of point 5.3 of the PN-C-89273:1997 standard should be stored at room temperature 18−22°C and relative humidity 50−70%.

**Accelerated aging with temperature factor**

The accelerated aging process was carried out in accordance with ASTM F1980-16 standard, using temperature as the aging factor [9, 10].

![Fig. 1. Accelerated aging of polymer](image-url)
The time to carry out the accelerated aging process was determined on the basis of the coefficient characteristics of the Arrhenius method as shown in Fig. 1. The Binder KMF 240 climatic chamber (Binder GmbH, Germany) was used, enabling the achievement of the temperature (20–80)±2°C and relative humidity < 20±5%. The storage life of the films was determined using the Arrhenius equation.

The following test conditions were established: temperature 65±2°C and relative humidity below 20%. The Arrhenius equation was used to determine the time intervals in the aging process:

$$AAF = \frac{Q^{10}}{10^{\frac{TAA-TRT}{10}}}$$

where:

- $TAA$ – accelerated aging temperature, °C
- $TRT$ – ambient temperature °C

It was assumed that one year of ageing would be achieved by incubating the sample for 18.5 days, two years for 37 days, 3 years – 55.5 days, 5 years – 92.5 days with air flow ≤ 1ml/s and atmospheric pressure of 860–1060 hPa.

### Accelerated aging with UV factor

The accelerated aging process was conducted on the basis of the guidelines specified in the following standards: PN-EN ISO 4892-1:2016-06, PN-EN ISO 4892-2:2013-06 and PN-EN ISO 105-B02:2014-11.

Time interval was 3 and 5 years. With the pre-set intensity of 600 W/m² within the range of wavelength 300–800 nm, the equivalent of natural radiation energy within month in the climatic zone of Central Europe in the UV aging chamber is achieved within 79 hours (the average annual sum of radiation dose for Poland is 3600 MJ/m²) [13–15]. Aging under an influence of UV radiation was carried out in a chamber for aging tests with a xenon arc lamp in an air-cooled xenotest Alpha+ (Atlas Material Testing Technology GmbH, Germany), which allows the following conditions to be achieved: wavelength $\lambda = 300–400$ nm, radiation intensity 160 W/m², temperature 38–40°C (black thermometer temperature 75°C) and humidity 40%, testing time was 200 h, during which the radiation dose was 109899 kJ/m² [15].

### Table 1. Physical parameters of the tested films

<table>
<thead>
<tr>
<th>Film</th>
<th>Mass per unit area g/m²</th>
<th>Thickness mm</th>
<th>Density g/cm³</th>
<th>Composition acc. to the manufacturer's declaration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>166±2</td>
<td>0.19±0.02</td>
<td>0.87±0.03</td>
<td>PET</td>
</tr>
<tr>
<td>Sample 1*</td>
<td>164±3</td>
<td>0.17±0.02</td>
<td>0.96±0.02</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 1**</td>
<td>156±4</td>
<td>0.16±0.02</td>
<td>0.97±0.02</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 2</td>
<td>173±6</td>
<td>0.17±0.02</td>
<td>1.04±0.03</td>
<td>PET</td>
</tr>
<tr>
<td>Sample 2*</td>
<td>180±6</td>
<td>0.19±0.02</td>
<td>1.00±0.05</td>
<td>PET</td>
</tr>
<tr>
<td>Sample 2**</td>
<td>168±1</td>
<td>0.17±0.02</td>
<td>0.99±0.01</td>
<td>PET</td>
</tr>
<tr>
<td>Sample 3</td>
<td>100±1</td>
<td>0.10±0.02</td>
<td>0.98±0.03</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 3*</td>
<td>107±3</td>
<td>0.11±0.02</td>
<td>1.00±0.05</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 3**</td>
<td>106±1</td>
<td>0.11±0.02</td>
<td>0.97±0.01</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 4</td>
<td>109±1</td>
<td>0.10±0.02</td>
<td>1.09±0.01</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 4*</td>
<td>117±1</td>
<td>0.10±0.02</td>
<td>1.17±0.01</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 4**</td>
<td>116±2</td>
<td>0.10±0.02</td>
<td>1.16±0.02</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 5</td>
<td>102±1</td>
<td>0.08±0.02</td>
<td>1.27±0.01</td>
<td>PES/PE/PP</td>
</tr>
<tr>
<td>Sample 5*</td>
<td>106±1</td>
<td>0.09±0.02</td>
<td>1.18±0.01</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 5**</td>
<td>108±2</td>
<td>0.09±0.02</td>
<td>1.20±0.03</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 5***</td>
<td>105±2</td>
<td>0.09±0.02</td>
<td>1.21±0.08</td>
<td>PP/PN</td>
</tr>
<tr>
<td>Sample 6</td>
<td>5.85±0.10</td>
<td>0.13±0.001</td>
<td>0.45±0.01</td>
<td>PET</td>
</tr>
<tr>
<td>Sample 6*</td>
<td>5.89±0.05</td>
<td>0.11±0.001</td>
<td>0.53±0.02</td>
<td>PET</td>
</tr>
<tr>
<td>Sample 6**</td>
<td>5.99±0.10</td>
<td>0.012±0.001</td>
<td>0.49±0.05</td>
<td>PET</td>
</tr>
<tr>
<td>Sample 7</td>
<td>51.6±0.1</td>
<td>0.59±0.003</td>
<td>0.87±0.02</td>
<td>PP</td>
</tr>
<tr>
<td>Sample 7*</td>
<td>54.6±0.1</td>
<td>0.59±0.003</td>
<td>0.93±0.01</td>
<td>PP</td>
</tr>
<tr>
<td>Sample 7**</td>
<td>54.0±0.9</td>
<td>0.66±0.021</td>
<td>0.77±0.01</td>
<td>PP</td>
</tr>
</tbody>
</table>
RESULTS AND DISCUSSION

The physical and mechanical properties of films are assessed in a laboratory using appropriately selected research methods. With the reservation that the more the property determination method is closer to actual conditions of use, the closer the assessment to the conditions of actual use.

The laboratory conditions also allow the simulation of accelerated aging in the temperature and UV radiation terms, reflecting the lifetime of the material.

With regard to the selected mechanical parameters, the graphic interpretation of the obtained test results is presented in Figs. 2–7.

Changes in the value of tensile strength in the longitudinal direction after aging for 3 and 5 years range from 0.5 to 41.5% (Fig. 2). The best result was obtained for sample 5 (PES/PE/PP). In this case, the tensile strength has practically not changed, which suggests the best resistance to thermal degradation. The greatest decrease in tensile strength was observed for sample 2 (PET), which proves the lowest resistance to thermal degradation.

The observed changes in the value of tensile strength in the crosswise direction after the aging periods of 3 and 5 years range from 0.5% to 37% (Fig. 3). Sample 6 (PET) showed the highest resistance to thermal aging and sample 2 (PET) the lowest. The differences in sensitivity to thermal aging of samples 6 and 2 (both PET films) indi-
cate a chemical modification of sample 6, which increased its resistance to aging.

As expected, the changes in elongation at break in both directions after the aging periods of 3 and 5 years are much greater than in the tensile strength, ranging from 1% to 63% (Figs. 4 and 5). Additionally, the sensitivity to thermal aging depends on the cutting direction of the samples (longitudinally or crosswise). In the longitudinal direction, sample 7 (PP) had the best thermal aging resistance, and sample 6 (PET) the worst. However, in the crosswise direction, the best resistance to thermal aging was recorded for sample 2 (PET), and the worst for sample 5 (PES/PE/PP).

It can be seen that the tested films are more sensitive to thermal degradation than UV radiation, as evidenced by changes in the tensile strength values after exposure to UV radiation, which do not exceed 14% (Fig. 6). The best resistance to UV was recorded for sample 4 (PP/PP), and the worst for sample 3 (PP/PA).

It can be seen from Fig. 7 that in both directions, the value of elongation at break after exposure to UV radiation increased only in the case of sample 4 (PP/PP), and for the
remaining samples it is reduced. This confirms the greater resistance of PP to UV than the other tested polymers.

**CONCLUSIONS**

Physical parameters (mass per unit area, thickness, density) after thermal and UV aging remain at the same level. Results do not differ between each other by more than 10%. The tested films are more sensitive to thermal degradation than UV radiation, as evidenced by changes in the tensile strength after exposure to UV radiation, which do not exceed 14%. The decrease in the value of tensile strength after the thermal aging does not exceed 40%. The lifetime of all the tested samples exceeds 5 years, in the case of none of the tested films, a 50% of decline decrease in the level of the measured parameters in relation to the initial value was achieved.

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