Comparison of efficacy of the intermittent pneumatic compression with a high- and low-pressure application in reducing the lower limbs phlebolymphedema

Introduction: The primary lymphedema and chronic venous insufficiency present an important medical problem, and effective physical therapeutic methods to treat this problem are still at the search phase. The aim of this study was to compare the efficacy of intermittent pneumatic compression (IPC) of a high- or low-pressure level in the treatment of primary phlebolymphedema of the lower limbs.

Methods: The study included 81 patients with chronic venous insufficiency and primary lymphedema of the lower limbs. Group A consisted of 28 patients who underwent a monthly antiedematous therapy including a manual lymphatic drainage, multilayer bandaging, and IPC with the output pressure of 120 mmHg. Group B consisted of 27 patients who underwent the same basic treatment as group A and IPC with the output pressure of 60 mmHg. Group C (control) consisted of 26 patients who underwent only a basic treatment – without IPC.

Results: After completion of the study, it was found that the greatest reduction of edema occurred in patients who underwent treatment with a pressure of 120 mmHg. The comparison of percentage reduction of edema showed a statistically significant advantage of the group A over groups B and C, both for the changes in the right (P=0.01) and the left limb (P=0.01). Results in patients undergoing intermittent compression of the lower pressure (60 mmHg) were similar to those obtained in the control group.

Conclusion: The IPC with the pressure of 120 mmHg inside the chambers effectively helps to reduce a phlebolymphedema. Furthermore, it appears that the treatments with a pressure of 60 mmHg are ineffective and their application becomes useless only in the antiedematous therapy.

Keywords: perometric measurement, intermittent pneumatic compression (IPC), phlebolymphedema, chamber pressure

Introduction
Lymphedema of the lower limbs, which occurs in patients with chronic venous insufficiency, is the result of disorders in tissue fluid, lymph, and venous blood circulation in a limb area, leading to a noticeable disparity in its appearance, reduced joint mobility, and frequent inflammation of the skin, blood vessels, and lymph nodes.¹-³

Currently, it is believed that the number of documented cases of lymphedema of the lower limbs could account for approximately 6 million people worldwide (4.6 million in the age group 40–60), in which 60% of patients suffer from chronic venous insufficiency.⁴ The only data collected in Poland, involving the most extensively studied population so far (40,096 patients contacting a General Practitioner [GP]), were
obtained in a multicenter cross-sectional study, whose results were published in 2003. The incidence of phlebolymphedema in patients diagnosed with chronic venous insufficiency in Poland was estimated to be 10%. Studies conducted in Western Europe show that the most common are telangiectasia and reticular veins as they occur in approximately 65% of women and >50% of men. In contrast, lower limbs varices affect 10%–20% of men and 25%–33% of women.

The results of epidemiological studies and papers referring to the financial costs of treating patients with phlebolymphedema clearly indicate the great need to seek minimally invasive, safe, and effective, but above all, low-cost therapeutic methods, and because of its chronic nature, also giving the opportunity to the patient to apply them at home.

Compression therapy is a method that exerts pressure on the limb from the outside, which increases the drainage from the veins and lymph, and this prevents the accumulation of fluid in the extravascular space.Compression therapy includes pneumatic massage device, multilayer limb bandaging, and compression garments with varying pressure. Intermittent pneumatic compression (IPC) is the physical method often used in sports and medicine, as evidenced by numerous bibliographic data from PubMed, Medline, and Physiotherapy Evidence Database (PEDro).

Despite a few clinical trials in lymphedema, a mutual consensus on the treatment parameters was not established. Some authors recommend the use of pressure from 40 mmHg to 60 mmHg, while others recommend significantly higher values from 80 mmHg to 130 mmHg.

The aim of this study was to compare the efficacy of IPC with a high- or low-pressure level in the treatment of primary phlebolymphedema of the lower limbs in patients with chronic venous insufficiency. It was evaluated whether the different level of output pressure in chambers during IPC procedures is a factor significantly influencing the effectiveness of antiedematous treatment in patients with chronic venous insufficiency and whether a high- or low-pressure value is more effective in edema reduction.

Materials and methods

Study design

This research project was approved by the Bioethics Committee of the Academy School of Physical Education in Katowice (number 5/2013, May 16, 2013). The research was conducted at the Limf-Med Hospital in Chorzow, in the period from July 1, 2013 to July 4, 2014.

The study included the patients with chronic venous insufficiency with unilateral or bilateral primary lymphedema of the lower limbs (lasting at least 3 months). Eligibility and clinical evaluation of patients were performed by a team consisting of a vascular surgeon and an angiologist, an internist, an oncologist, and a physiotherapist.

Patients were assigned to one of the three comparison groups, A, B, and C. The basis for inclusion of patients was a vascular Doppler examination with the use of the Doppler Duplex Hitachi EUB 5500 device. The study included an assessment of venous reflux in superficial and deep system and in the area of perforators. According to the standards, the venous reflux is present with a valve insufficiency of >0.5 seconds at the compression trial. The patients were also subjected to specification according to the Clinical Etiology Anatomy Pathophysiology (CEAP) classification of chronic venous insufficiency. The inclusion criteria also included positive Stemmer’s sign (Figure 1) and 20-second compression attempt of the skin (Figure 2). The severity of phlebolymphedema was evaluated based on two clinical classifications for lower limbs by Olszewski and Brunner. In the case of women to be eligible for treatment, each treatment was adjusted to the beginning of the follicular phase of the menstrual cycle and initiated only after menstruation. Exclusion criteria included patients with active thromboembolic disease, diabetes, peripheral arterial disease (ankle-brachial index <0.8), rheumatoid arthritis, parasitosis, generalized scleroderma, collagenoses, dermatitis, psoriasis, and advanced coronary artery disease (III and IV in New York Heart Association [NYHA] classification) – edema of cardiogenic origin, patients with polyneuropathy of the lower limbs, as well as patients with pacemakers and patients taking steroid and/or diuretic medications. The study also excluded patients with edema of renal or hepatic nature and patients with postoperative

Figure 1 Characteristic changes (infiltration of lymph) within the second toe – a massive Stemmer’s sign.
lymphedema (secondary) and cancer. Patients with irregular menstrual cycles (6-month observation before entering the study) in premenopausal period were also excluded.

Characteristics of the study population

Prior to the study, participants were briefed on the objective of the study and given the opportunity to refuse further participation at any point. All participants gave their informed consent to be included in the study (statement confirmation) and consent was also given by the patient for the use of the photo in Figure 6. The study was conducted in accordance with the principle of intention to treat (patients excluded from the research at subsequent stages for various reasons were still provided with a basic lymphatic care).

The group of 81 patients eligible for the treatment was assigned to three comparative groups. A random selection for particular groups was carried out continuously throughout the duration of the study, that is, each new patient was subjected to randomization by computer random number generator using a Monte Carlo method, and based on the result of the draw, the patient was assigned to a given control group according to the CONSORT guidelines (Figure 3).

The characteristics of the study population within each group are presented in Tables 1–4. The groups were homogeneous taking into account all the factors characteristic for the patients, as well as the initial size and the clinical stage of chronic venous and lymphatic edema of the lower limbs.

Treatment procedures

All patients were treated on an outpatient basis in a lymphatic clinic, where they received a 4-week comprehensive antiedema therapy (manual lymphatic drainage and multilayer bandaging) and pharmacotherapy (phlebotropic drug – Diosmin 500 mg two tablets administered per day).

The manual lymphatic drainage was performed by the physical therapist starting from the proximal preparation (ie, gentle strokes on the so-called venous angles – skin area over the subclavian veins), then the skin layers over the cisterna chyli (a saccular area of dilatation in the lymphatic channels as the beginning of the thoracic lymph duct) connected with the diaphragm breathing exercises, and finally segmental massage of the entire upper limb. The massage was finished with a proximal preparation technique. Lymphatic massage took 50 minutes and was performed once a day, three times a week (Monday, Wednesday, and Friday).

After performing the manual lymphatic drainage (before placing a limb low or resting), the compression therapy was applied – multilayer bandaging with an external pressure of 40–50 mmHg. The first layer was an orthopedic sleeve Tubula (Matopat Bella-Handelv LLC, Torun, Kujavy, Poland) applied directly onto the surface of the limb. Then the elastic Matolast support bandage (Matopat Bella-Handelv LLC) was used for accurate and precise overlay on the toes. The next layer was a RoltaSoft cotton (Paul Hartmann LLC, Pabianice, Lodzkie, Poland), which starts from the distal part and was gradually applied to the entire limb. The outer layer consisted of evenly placed short-stretch bandage (Paul Hartmann LLC), which was applied for 24 hours. In cases of bilateral edema, bandaging was performed on both limbs in accordance with the standards to the level of the knee joint and not to the height of the groin as in the case of bandaging a single limb with phlebolymphedema.27

In addition, the IPC treatment was performed in group A and B patients before manual lymphatic drainage treatments with the use of the 12-chamber apparatus Lymphatron DL1200 (Technomex LLC, Gliwice, Upper Silesia, Poland; Figures 4 and 5). In group A, the external pressure used was as high as 120 mmHg, and in group B, it was 60 mmHg. A single treatment took 45 minutes. The procedure consisted in sequentially filling the chambers of the sleeve starting from the circumference going to closer sections of a limb. A single-chamber filling time was 3 seconds (when the air was pumped to the next chamber, the previous ones did not deflate, which means that they maintained full pressure). In cases of unilateral edema, a single compression sleeve was used, and in patients with bilateral edema, a double sleeve was used. The treatment was conducted once a day, three times a week (Monday, Wednesday, and Friday). Manual lymphatic drainage treatments and multilayer bandaging were performed in all patients from groups A, B, and C by the same therapist.
Methods and measuring error

Lower limb volume measurements were performed by the same technician in all three groups of patients before and after a monthly therapy.

To assess the volume of a limb, an optoelectronic Perometer 400 T was used (Pero-System Messgeräte GmbH, Wuppertal, North Rhine-Westphalia, Germany; Figure 6), which was connected to a personal computer. This method allowed to estimate the volume of a limb with a measuring error of 0.5% (manufacturer’s data). The assessment technique was based on a special ring, equipped with a system of 378 Lighting Emitting Diode (LED) diodes emitting the infrared radiation. Within the ring, there were also the optical sensors that receive electromagnetic stimuli. During the course of measurement, the limb was located inside the ring on the diode-sensor lines. The registered light pulses on the detectors were turned into electronic signals. The ring was moved during measurement to cover the entire limb. Computer software allowed for the analysis of volume and circumference at any stage of measurement.

For the purpose of this research, we made our own estimates of measurement error of Perometer 400T. On the basis of ten patients who were randomly selected for the study, each of them was subjected to 20 consecutive measurements of lower limb with edema (a total of 200 measurements).
Table 1 General characteristics of the study population in groups A, B, and C

<table>
<thead>
<tr>
<th></th>
<th>Group A</th>
<th>Group B</th>
<th>Group C</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients</td>
<td>24</td>
<td>23</td>
<td>26</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>Sex</td>
<td>15F/9M</td>
<td>13F/10M</td>
<td>16F/10M</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>Number of smokers</td>
<td>13</td>
<td>12</td>
<td>13</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>Age (years)</td>
<td>Average</td>
<td>49.5</td>
<td>51.5</td>
<td>52.2</td>
</tr>
<tr>
<td></td>
<td>Min–max</td>
<td>42.3–62.2</td>
<td>44.6–60.3</td>
<td>45.6–63.4</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>Average</td>
<td>26.4</td>
<td>27.7</td>
<td>27.8</td>
</tr>
<tr>
<td></td>
<td>Min–max</td>
<td>22.3–32.5</td>
<td>19.6–32.2</td>
<td>20.2–32.4</td>
</tr>
<tr>
<td>Serum creatinine (mg/dL)</td>
<td>Average</td>
<td>1.5</td>
<td>1.3</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td>Min–max</td>
<td>0.8–1.4</td>
<td>1.1–1.8</td>
<td>1.1–1.9</td>
</tr>
</tbody>
</table>

Notes: Group A underwent a monthly antiedematous therapy including a manual lymphatic drainage, multilayer bandaging, and IPC with the output pressure of 60 mmHg. Group B underwent the same basic treatment as group A and IPC with the output pressure of 120 mmHg. Group C (control) underwent only a basic treatment – without IPC.

Abbreviations: BMI, body mass index; F, female; L, left limb; max, maximum; M, male; min, minimum; R, right limb; SD, standard deviation; IPC, intermittent pneumatic compression.

The absolute error ($\Delta X$) of the measurement of the total lower limb volume was calculated using the formula:

$$\Delta X = X - X_0,$$  \hspace{1cm} (1)

where $X$ is the value measured in the next trial and $X_0$ is the correct value (mean) calculated after 20 trials.

Then, the relative error was estimated ($\delta X$) from the formula:

$$\delta X = \frac{\Delta X}{X_0},$$  \hspace{1cm} (2)

where $\Delta X$ is the absolute error and $X$ is the value measured at a given trial.

Table 2 Characteristics of chronic venous insufficiency in given groups

<table>
<thead>
<tr>
<th>CEAP scale*</th>
<th>Group A</th>
<th>Group B</th>
<th>Group C</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1 E, A <em>2,3</em> R</td>
<td>10R/10L</td>
<td>9R/11L</td>
<td>10R/11L</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>C1 E, A <em>2,3</em> R</td>
<td>8R/7L</td>
<td>8R/7L</td>
<td>9R/8L</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>C1 E, A <em>2,3</em> R</td>
<td>6R/7L</td>
<td>6R/5L</td>
<td>7R/7L</td>
<td>&gt;0.05</td>
</tr>
</tbody>
</table>

Notes: C, CEAP scale; E, primary failure; C, primary failure; edema, and trophic changes in the skin and subcutaneous tissue; the presence of reflex above and below the knee in saphenous vein and femoral vein; and C, primary failure, edema and trophic changes in the skin and subcutaneous tissue, the presence of reflex above and below the knee in saphenous, femoral, and popliteal vein and shank perforators. Group A underwent a monthly antiedematous therapy including a manual lymphatic drainage, multilayer bandaging, and IPC with the output pressure of 120 mmHg. Group B underwent the same basic treatment as group A and IPC with the output pressure of 60 mmHg. Group C (control) underwent only a basic treatment – without IPC.

Abbreviations: R, right limb; L, left limb; IPC, intermittent pneumatic compression; CEAP, clinical etiology anatomy pathology.

Then, the mean percentage error (relative error expressed in percentage) of all 20 measurements for each limb was calculated. 28

The final measurement error by the authors’ calculations (based on the error results of ten lower limbs) was higher than that specified by the manufacturer (average = 1.26%, and standard deviation [SD] = 1.03%). The results of the measurement error were obtained for three right and seven left lower limbs (mean limb volume amounted to 15,532.44 cm³, and 5,092.31 cm³ SD).

The tests with perimeter measured the changes of real values (absolute) in the total volume of the limb and its sections – the volume of the foot, shank, and thigh – and additionally percentage changes for edema reduction were calculated using the following formula:

$$dP\% = \frac{(P_f - P_i) \times 100\%}{P_i},$$  \hspace{1cm} (3)

where $dP\%$ is the percentage change in a given parameter, $P_i$ the initial real value of a given parameter (measured before treatment), and $P_f$ the final real value of a given parameter (measured after treatment). 28

Table 3 Patients’ characteristics in terms of edema according to categories by Olszewski

<table>
<thead>
<tr>
<th>Edema severity according to Olszewski’s*</th>
<th>Group A</th>
<th>Group B</th>
<th>Group C</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>IIa (number of patients)</td>
<td>5R/3L</td>
<td>4R/4L</td>
<td>4R/3L</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>IVa (number of patients)</td>
<td>10R/9L</td>
<td>10R/8L</td>
<td>12R/10L</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>IVc (number of patients)</td>
<td>2R/1L</td>
<td>2R/1P</td>
<td>3R/2L</td>
<td>&gt;0.05</td>
</tr>
</tbody>
</table>

Notes: *Type IIa, pitting edema in the foot and shank; type IVa, pitting edema in the entire limb; and type IVc, edema with hyperkeratosis, fibrosis, and leakage of lymph in the entire limb. Group A underwent a monthly antiedematous therapy including manual lymphatic drainage, multilayer bandaging, and IPC with the output pressure of 120 mmHg. Group B underwent the same basic treatment as group A and IPC with the output pressure of 60 mmHg. Group C (control) underwent only a basic treatment – without IPC.

Abbreviations: R, right limb; L, left limb; IPC, intermittent pneumatic compression.
Table 4 Patients’ characteristics in terms of edema according to categories by Brunner

<table>
<thead>
<tr>
<th>Edema severity according to Brunner (number of patients)</th>
<th>Group A</th>
<th>Group B</th>
<th>Group C</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>II 2R/1L</td>
<td>2R/1L</td>
<td>3R/2L</td>
<td>&gt;0.05</td>
<td></td>
</tr>
<tr>
<td>III 9R/7L</td>
<td>8R/7L</td>
<td>9R/6L</td>
<td>&gt;0.05</td>
<td></td>
</tr>
<tr>
<td>IV 2R/1L</td>
<td>2R/1L</td>
<td>3R/2L</td>
<td>&gt;0.05</td>
<td></td>
</tr>
<tr>
<td>V 1R/1L</td>
<td>1R/1L</td>
<td>1R/1L</td>
<td>&gt;0.05</td>
<td></td>
</tr>
</tbody>
</table>

Notes: ‘Grade II, all-day edema resolves spontaneously after the night, with positive Stemmer’s sign; grade III, constant edema does not pit and does not significantly reduce upon limb elevation; grade IV, permanent edema distorting a limb, complicated as associated with irreversible skin changes, ie, fibrosis and papillae, as a result of inflammation of the skin; grade V, elephantiasis, massive edema distorting a leg with skin thickening, and changes in muscles – dystrophy, disabling the function of a limb. Group A underwent a monthly antiedematous therapy including a manual lymphatic drainage, multilayer bandaging, and IPC with the output pressure of 120 mmHg. Group B underwent the same basic treatment as group A and IPC with the output pressure of 60 mmHg. Group C (control) underwent only a basic treatment – without IPC.

Abbreviations: R, right limb; L, left limb; IPC, intermittent pneumatic compression.

Statistical analysis
For the analysis of the parameters studied in the paper, the STATISTICA software version 10.0 (StatSoft Poland, Dell Inc., USA) was used. The uniformity of distribution of the patients’ characteristics in groups A, B, and C was tested with the chi-squared test version of the highest reliability ($\chi^2$) and with the Kruskal–Wallis homogeneity test. Statistically significant differences were considered when $P<0.05$. The nonparametric matched-pair Wilcoxon test was used to check whether a given parameter changed statistically significantly after the treatment compared to initial value ($P<0.05$) within group (dependent variables). In contrast, the nonparametric test for independent variables – Kruskal–Wallis analysis of variance (ANOVA) – examined the relationship between percentage changes in groups A, B, and C. However, the post hoc Tukey’s test for multiple comparisons was used to find exact dependence resulting from analysis of variables between groups, differences in the changes between the groups in the limb volume, and edema values. Differences were statistically significant when $P<0.05$.

Results
The analysis of dependent variables with the use of Wilcoxon matched pair test found a statistically significant decrease in the volume of right lower limb as compared to the initial parameters in all comparison groups (Table 5).

The Kruskal–Wallis ANOVA indicates a statistically significant difference between the groups $P(A, B, C) = 0.00000$. Post hoc test indicated a significant advantage of group A in terms of a percentage reduction in the volume of the affected lower limb in relation to other comparison groups. By contrast, there was no difference observed between groups B and C: $P(A vs B) = 0.000002$, $P(A vs C) = 0.000000$, and $P(B vs C) = 0.7052$. The detailed results of the percentage changes in the volume of right lower limb are shown in Figure 7.

Likewise, a statistically significant reduction in the volume of the left lower limb in relation to the initial parameters was observed in each group (Table 6). It was also reported that there was a statistically significant difference between the groups $P(A, B, C) = 0.0000$. Post hoc test indicated a significant advantage of group A in terms of a percentage decrease in the volume of the left lower limb.
reduction in the volume of the affected left lower limb in relation to other comparison groups. By contrast, there was no difference observed between groups B and C: $P(A \text{ vs } B) = 0.000000$, $P(A \text{ vs } C) = 0.000002$, and $P(B \text{ vs } C) = 0.7093$. The detailed results of the percentage changes in the volume of left lower limb are shown in Figure 8.

**Discussion**

The worldwide consensus on the IPC parameters (high or low pressure in chambers) is not established. Olszewski believes that the external pressure exerted on the surface of a limb in order to effectively reduce accumulated lymph should be several times greater than the pressure inside the vessel acting directly on the outflow. He found, with the use of implanted cannula, that if the mean physiological pressure in the superficial lymphatic vessel in the lower limb is 10–12 mmHg, then the external pressure (on skin) is 30–40 mmHg. Therefore, only greater external pressure can stimulate lymph drainage. For example, assuming a mechanical external compression equals 60 mmHg, the pressure acting inside the lymph vessel will amount to only approximately 15–20 mmHg (and in accordance with the basic laws of physics by Laplace and Poiseuille, the flow can be initiated only at this value). In contrast, when the applied pressure is 100 mmHg, then the intravascular pressure may increase to 30–40 mmHg. Olszewski calculated that under any additional pressure of 2 mmHg, the lymph flow increases by a further 50 mL and recommends the use of the external pressure from 80 mmHg to even 150 mmHg, and the treatment time should be 45–60 minutes.

On the other hand, Ponikowska et al argue that the level of pneumatic compression pressure should not exceed

### Table 5 Between-group comparisons of changes in right lower limb volume (cm$^3$)

<table>
<thead>
<tr>
<th>Group</th>
<th>Before treatment</th>
<th>After treatment</th>
<th>$P$-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group A</td>
<td>Average</td>
<td>14,423.34</td>
<td>8,755.23</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>4,781.12</td>
<td>7,878.77</td>
</tr>
<tr>
<td>Group B</td>
<td>Average</td>
<td>14,562.02</td>
<td>12,511.47</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>4,454.24</td>
<td>8,954.43</td>
</tr>
<tr>
<td>Group C</td>
<td>Average</td>
<td>14,078.22</td>
<td>12,305.67</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>4,303.88</td>
<td>9,887.98</td>
</tr>
</tbody>
</table>

**Notes:** Group A underwent a monthly antiedematous therapy including a manual lymphatic drainage, multilayer bandaging, and IPC with the output pressure of 120 mmHg. Group B underwent the same basic treatment as group A and IPC with the output pressure of 60 mmHg. Group C (control) underwent only a basic treatment – without IPC.

**Abbreviation:** SD, standard deviation; IPC, intermittent pneumatic compression.

### Table 6 Between-group comparisons of changes in left lower limb volume (cm$^3$)

<table>
<thead>
<tr>
<th>Group</th>
<th>Before treatment</th>
<th>After treatment</th>
<th>$P$-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group A</td>
<td>Average</td>
<td>13,902.11</td>
<td>8,734.23</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>4,423.33</td>
<td>6,956.44</td>
</tr>
<tr>
<td>Group B</td>
<td>Average</td>
<td>14,103.23</td>
<td>12,567.78</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>4,223.25</td>
<td>9,123.84</td>
</tr>
<tr>
<td>Group C</td>
<td>Average</td>
<td>13,944.21</td>
<td>12,270.67</td>
</tr>
<tr>
<td></td>
<td>SD</td>
<td>3,993.84</td>
<td>9,563.86</td>
</tr>
</tbody>
</table>

**Notes:** Group A underwent a monthly antiedematous therapy including a manual lymphatic drainage, multilayer bandaging, and IPC with the output pressure of 120 mmHg. Group B underwent the same basic treatment as group A and IPC with the output pressure of 60 mmHg. Group C (control) underwent only a basic treatment – without IPC.

**Abbreviation:** SD, standard deviation; IPC, intermittent pneumatic compression.
In the treatment of edema in patients with venous insufficiency, and the treatment may be not tolerated because of excessive pressure on the pain receptors in the skin.

Similar guidelines are also recommended by Woźniewski and Kołodziej, explaining that the pressure in the pneumatic compression >60 mmHg should not be applied in patients with lymphedema of the limbs (also of venous component) because higher pressure can be painful and may damage the skin vessels (lymphatic precollectors in the skin).

This paper is the first attempt, in the available literature, to compare two different (relatively low and high), but frequently used in everyday therapeutic practice (and sometimes even despite conflicting physical basis and the lack of reliable scientific foundations and reliable clinical trials) pressures in the IPC treatment within one experiment and based on a uniform study material, including patients with specially selected edema with mixed component, that is, phlebolymphedema. The innovation was also the use of accurate and noninvasive measurement of lower limb edema with the use of an innovative perometric test. So far, researchers have used only the methods with high measurement error, namely, a metric technique using a metric tape, to calculate the circumference of the limb volume from the formula for a truncated cone or by means of water displacement (hydrostatic method). The optoelectronic device Perometer 400 T, used in our study, was the first trial in scientific research in the field of phlebolymphedema of the lower limbs.

Therefore, the obtained results can only be referred to individual reports in the literature on pneumatic compression therapy used in the treatment of edema in patients with chronic venous insufficiency of the lower limbs or slightly more numerous publications in the field represented by the therapy of the so-called secondary lymphedema, where the most research material include patients after gynecological and oncological surgeries with resection of the inguinal and/or retroperitoneal lymph nodes.

The results obtained by Zaleska et al really correspond to our experience and seem to justify the application of high pressure inside the chambers during IPC, both in cases of mixed phlebolymphedema and postoperative lymphatic edema resulting from inguinal node resection after hysterectomy. The study used a multichamber compression sleeve, supporting the opinion that its effectiveness during treatment is higher than a single-chamber sleeve. A single-chamber filling time was different – relatively short and lasted 3 seconds and not 50 seconds (in the future, it is worth considering to conduct a comparative analysis of the clinical effectiveness of 50-second duration of compression in the case of phlebolymphedema). Besides, still three patients experienced discomfort and pain sensations during intermittent compression with a pressure of 120 mmHg, which led to termination of the treatment. In group B, where applied pressure value was 60 mmHg, there were two such cases. Our studies do not fully support the thesis that this therapy (especially high pressure) is always well tolerated by patients.

Abu-Own et al also advocated the use of multichamber compression. The study was performed in 17 healthy volunteers, where two types of devices for IPC were applied (multichamber compression exerting the compression of 40–55 mmHg and a single-chamber device exerting pressure all over the shank at the same time under a pressure of 60 mmHg) for a period of 10 minutes. The result was a significantly greater reduction in volume after each pressure when using the multichamber compression.

Similar conclusions on the application of higher pressure parameters in the treatment of lymphedema were drawn by Modaghegh and Soltan, stating that IPC is an extremely effective physical treatment (16-hour treatment and pressure value of 80–120 mmHg). It was found that the average reduction of edema in the affected limb was as high as approximately 75%.

The available literature also presents critical works on the use of high pressure during IPC in cases of primary and secondary lymphatic edema of lower limbs. However, it must be critically assessed that most of these works lack comparison groups and blindness, and the studies were based on a small number of patients.
The literature does not devote much place for studies in basic science. Certainly, a deeper understanding of all the mechanisms that occur under the influence of IPC within the lymphatic vessels and lower limb venous system would facilitate clinical activity. An interesting aspect of the testing is the analysis of thermographic images taken with infrared camera in patients with venous edema and trophic disorders of the skin (open sores). The research shows that the applied compression with pressure >60 mmHg increases the temperature of the lower leg and the temperature around the medial malleolus of treated patients (which is justified by stimulation of microcirculation). The lower pressure values did not cause this effect.23

In our work, we did not evaluate the financial costs of IPC; however, one has to agree with the fact that this method – apart from a one-off expense for the purchase of equipment – allows to perform physiotherapy treatments at home and does not require complicated service. Besides, it is available and relatively cheap (as for medical devices).

Limitations of study
There are limitations to this study, for example, the small number of subjects and lack of the follow-up results. Another potential limitation of our study is the lack of a pain relief and quality of life observation – larger aspects than only edema therapy. We acknowledge the need to continue this research in larger numbers of subjects, thus, according to statistical estimation, we would like to clarify that the population over 35–40 in each group is needed for further analysis of normal distribution and to use the parametric tests. At this moment, it was estimated that a power of test is 0.86 compared to parametric statistics conducted on large population with Gaussian decay (30 subjects in one group). At the same time, we emphasize the pilot character of this study, which describes the insufficient utility of IPC with low pressure for lower leg phlebolymphedema. Our results should be confirmed on bigger population (perhaps with sham IPC – no pressure in chambers) especially because of lack of the sample size estimation.

Conclusion
The output pressure in chambers during IPC treatments is a factor that significantly affects the effectiveness of edema treatment, in patients with chronic venous insufficiency. IPC with a pressure inside the chambers of 120 mmHg significantly helps to reduce phlebolymphedema. In contrast, it appears that treatments at a pressure of 60 mmHg are ineffective and their use only in antiedematous therapy becomes pointless (in the scope of the study).

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References


