BTL-4000 Topline
Ultrasound Therapy

USER'S GUIDE
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Continuing education is a very important aspect of healthcare delivery. Many excellent resources are available today to expand a user's knowledge about many aspects of ultrasound therapy. BTL recommends a thorough review of this guide prior to operating the equipment.

1.1 Mechanism of Action

Mechanical waves of frequency higher than 20,000 Hz are called ultrasound. Application of ultrasound does not generate any electric current in tissues, and thus it is classified as mechanotherapy. The frequency usually used in physiotherapy is 0.8 – 3 MHz.

When the air gap between the emitter head and the body surface is eliminated, the emitter head vibrations are transmitted into the tissue and propagate to depth in the form of longitudinal waves. All cells in the path of the ultrasonic beam begin to oscillate. This causes micro-massage followed by transformation of gel into sol (jelly structures become liquid), transformation of mechanical energy into thermal, and deep warming of tissues.

The amount of generated heat depends on the amount of absorbed energy. Other effects of ultrasound (degassing of solutions, formation of cavities in liquids, and local alkalization) are negligible with the ultrasound doses and intensities used in physical therapy.

In order to avoid side effects, it is important to realize that molecular oscillation occurs not only in the path of the ultrasonic beam, but also in places that are distant from the area of application as a result of transmission by body fluids. This may lead to restoration of former epistaxis or acceleration of menstruation.

The features of the ultrasound beam and its distance from the emitter head determine the ultrasound field as being either close or distant.

**Close ultrasound field** is characterized by low beam divergence and large intensity variations due to interferential effects. The length of the close field is directly proportional to the effective radiation area (ERA – see below) of the emitter head and inversely proportional to frequency. For example, the length of a close field with 4 cm² ERA emitter head and 1 MHz frequency is approximately 10 cm, and for an emitter head with 1 cm² ERA and 1 MHz frequency it is approximately 2 cm.

**Distant ultrasonic field** is characterized by increasing beam divergence, gradual decrease in intensity, and almost no interferential effects.

Therapeutic effects take place mainly in the close field. The ultrasonic beam in the close field has significant interferential effects (interference of applied and reflected waves) - both constructive and destructive. It results in a non-homogenous ultrasonic beam where peak levels of intensity (local increase in intensity caused by constructive interference) may be many times higher than the pre-set value.

**Beam Non-uniformity Ratio (BNR)** states how many times the peaks of intensity exceed the pre-set values. This value characterizes an ultrasound head with a given frequency. The BNR value of a good quality ultrasound head is lower than 5. This means that if the pre-set intensity of the unit is 1 W/cm², the intensity in any part of the ultrasonic beam is not higher than 5 W/cm².

The BNR of older ultrasound heads and some newer ones (some manufacturers do not mention the BNR value) is often 20 or even more!

The Effective Radiating Area (ERA) is always smaller than the actual surface area of the emitter head (the ERA is determined by the size of the piezoelectric crystal or ceramic tablet that generates ultrasound by oscillating).
The ultrasound dose (power emitted to a surface area) is, therefore, related to the ERA and not to the actual surface area of the emitter head.

The phenomena of ultrasound refraction and reflection are caused by ultrasound wave transmission from one tissue into another and by different transmission speeds in these tissues. When applying ultrasound, it is necessary to eliminate the air gap between the emitter head and the skin. Therefore, modern ultrasound heads have a built-in optical and/or acoustic check of insufficient contact, possibly with automatic termination of application time countdown.

Due to interference in the close ultrasound field (it reaches the highest level at the site of the soft tissue/bone boundary - up to 35%), ultrasonic beam power increases (constructive interference) or decreases (destructive interference).

In order to avoid tissue lesions at peak levels of intensity, it is necessary to move the emitter head continuously.

As a result of reflection and constructive interference, local increase in intensity and temperature may occur, particularly at the periosteum/bone boundary. This increase can lead to periosteum pain during ultrasound application. If this occurs, the intensity must be immediately lowered.

Ultrasound is primarily absorbed in deeper tissues. Since these tissues usually do not include thermoreceptors, it is impossible to perceive local rise in temperature. The patient feels pain only if the local temperature exceeds 45°C and nociceptive receptors are irritated. Most authors agree that a short-term rise in local temperature to 45°C is not dangerous.

In the area of classic inflammation (oedema, erythema, local rise in temperature, pain or dysfunction) additional heat production is contraindicated, and thus only pulsed ultrasound (athermic) can be applied if necessary.

During the peracute phase of post-traumatic conditions (up to 24-36 hours) pulsed ultrasound application is contraindicated (vibrations hinder capillary proliferation and may cause delayed bleeding).

Local rise in temperature and micro-massage have several physiological effects:

- **Improvement of local circulation and thus also metabolism.** Rise in temperature enhances vasodilatation (more evident in continuous ultrasound).
- **Increase in capillary permeability and increased resorption of extravasation fluid.**
- **Improvement of local circulation and decrease in orthosympathetic activity resulting in significant muscle relaxation.**
- **Decrease in local ischaemia pain.**
- **Transformation of gel into sol** (due to transformation of fibrinogen into fibrin, haematomas and oedemas change into gel; ultrasound dissolves this gel and speeds up resorption). As the transformation of fibrinogen into fibrin is the basis of the healing process (scar formation), it is not advisable to apply ultrasound in peracute post-traumatic conditions.
- **Improvement of tissue regeneration capabilities as a result of the above-mentioned effects.**
Ultrasound also has several non-therapeutic effects that can have negative impacts, such as:

- **Tissue lesion** - Mechanical and/or thermic tissue lesion can occur when the intensity is too high. Especially sensitive is the nervous system (peripheral nerve) situated right on the bone (interference!) below the surface (close field!). Impulse transmission speed in the corresponding nerve decelerates, then occurs total (reversible) impulse transmission blockage and finally irreversible disintegration of the neuron (myelin coats are preserved). Therefore, be extremely cautious when applying ultrasound e.g. on paravertebral muscles after laminectomy when nerve structures lose their natural bone protection. Similarly, bone projections located just under the skin (ankles, epicondyles, spondyle spines, etc.) are also sensitive.

- Leukocyte mobility impairment – this can be minimized by sufficient movement of the emitter head.

- Other effects (mainly caused by overdosage) are: decrease in glycaemia, increased fatigue, nervousness, changes in appetite, constipation, increased tendency to catch colds.

### 1.2 METHODS OF APPLICATION

#### 1.2.1 ACCORDING TO APPLICATION AREA AND MOVEMENT OF THE EMITTER HEAD

**Static application** - The head is fixed to the treated area by a special holder and it is not moved. Due to the above-mentioned adverse effects, it is the least suitable form of application.

**Semi-static application** - is used when the application area corresponds to the ERA of the emitter head. The therapist continuously moves the emitter head in a spiral along the perimeter of an imaginary circle.

**Dynamic application** - the application area is bigger than the ERA of the emitter head. The therapist moves the emitter head in spirals in the treatment area. The application time is prolonged proportionally according to how many times the application area is bigger than the ERA of the head.

#### 1.2.2 ACCORDING TO APPLICATION AREA

**Local application** - Ultrasound is applied to the affected area. It is the most common manner of application, particularly suitable for local muscle spasms, chronic post-traumatic edemas, etc.

**Segmental application** - Ultrasound is applied to the outflows of nerve radices of the affected area (e.g. Sudeck syndrome, Morbus Reynaud, etc). The application is paravertebral and homolateral; in the area of C5 - Th1 for pathologies of upper limbs and in the L3 - S1 area for lower limbs.

**Neural application** - is based on the effect of decrease in impulse transmission speed in the peripheral nerve where ultrasound is applied. As the dividing line between the lowering of conductivity and irreversible nerve damage (asymptomatic) is very fine, this method of application is considered dangerous, and it is used only exceptionally (e.g., phantom pain).

**Radicular application** - Ultrasound is applied subsequently to the corresponding spinal root and manifested Head zone. For application above the spinal root, the same risks and limitations apply as for neural application.

#### 1.2.3 ACCORDING TO CONTACT BETWEEN EMITTER HEAD AND BODY SURFACE

**Direct contact** - is provided by a contact medium (ultrasound gel). This is a common manner of ultrasound application, and it is not necessary to mention it in ultrasound therapy prescription. For combined therapy (ultrasound + electrotherapy) it is recommended to write "conductive gel" in the prescription because some gels do not conduct electricity. The BTL ultrasound gel is conductive; if this gel is not available, you can use ECG gel or another water-based gel instead. However, never use paraffin oil, because it is non-conductive and could damage the head.
Subaqual application. This method has a range of advantages: It uses mainly the distant ultrasound field, where interference does not occur. Furthermore, there is no need to press the emitter head against the skin in order to maintain sufficient contact (this pressure is unpleasant or even painful for post-traumatic conditions). In addition, this application is not limited by uneven surfaces, and ultrasound can be easily applied to these surfaces, such as to interphalangeal joints.

Disadvantages of subaqual application include difficult handling of the special porcelain bath, limitation of ultrasound application to acral body parts, and the risk of the therapist getting hand lesions if the hand is put in water. (It is strictly prohibited due to reflection and interference of ultrasonic waves against the walls of the porcelain bath!) Some emitter heads (even newer) are claimed to be water resistant, but if they do not have a holder, they do not provide safe subaqual ultrasound application.

Ultrasound can also be applied via a thin-walled rubber bag (surgical gloves, condom) filled with boiled water. However, when using this method, it is necessary to eliminate the air gap between the bag and skin (use gel) and between the emitter head and the bag. This method is often perceived as very time-consuming.

1.3 ULTRASOUND FREQUENCY

Older ultrasound units employ a fixed frequency, usually 0.8 – 1 MHz. Newer ones are multi-frequency units, the selected frequency being determined by the target tissue; 1 MHz frequency is used for deep tissues and 3 MHz frequency for superficial tissues.

1.4 MODES OF OPERATION

a) continuous - is characterized by heat generated deeply in tissues. It is contraindicated in inflamed areas and anywhere else where local warming is undesirable.

b) pulsed - together with shortening of pulse length there occurs decrease in the so-called Duty factor. As a result, the thermic effect is suppressed and when the Duty factor is below 12.5% (1:8), an athermic effect can be expected.

Duty factor (DF) can be set in pulsed mode of operation. Its value states for how many percent of the duration the ultrasound signal is generated. When setting therapy parameters of BTL units, the ratio of ultrasound signal duration to the period length is stated in brackets. This ratio is used in the parameters of recommended ultrasound therapies of BTL units. In case of continuous setting of the parameter, only the percentage expression is used.

Example:

Duty factor 25% (1:4) means that 25% (1/4) of the period is ultrasound and the rest of the period is pause.

The same example, more detailed:

With a pulse frequency of 100 Hz, the period is 10ms long. With DF set to 25% (1:4), pulse duration is 2.5 ms and pause is 7.5 ms.

1.5 Emitter Head Size

The size of the emitter head is determined by the Effective Radiating Area (ERA): 1 cm² (small head) and 4 cm² (medium head, or large head according to some manufacturers). The choice of the head depends on the size of the application area. Ultrasound application to large areas using a small head is too time-consuming, difficult for the therapist, and the dose is not applied homogeneously.

Small heads with ERA about 1 cm² are used for trigger points, scars or for small and uneven surfaces where with an emitter head of standard size the ultrasound would have to be applied subaqualy.
1.6 APPLICATION TIME

Application time varies and largely depends on the stage of the disease. For acute conditions, it is 3 minutes at the beginning; for chronic conditions, it is usually 5 minutes at the beginning, and then it is prolonged using the positive step method. Application time for most therapies does not exceed 10 minutes. If the application area is x-times larger than the ERA of the head, application time has to be x-times prolonged and dynamic application is used.

1.7 INTENSITY

For acute conditions, start with an intensity of 0.5 W/cm² (in some cases even lower: e.g., in case of trismus, 0.2 W/cm² on hypertonic fibres of m. temporalis. The muscle is situated on the bone, just below the surface. Since the close ultrasound field is used, apply very low intensity).

For chronic conditions, an intensity of 0.8 - 1.0 W/cm² is used initially and it is increased according to the patient's reaction (the positive step method). The ultrasound intensity must not exceed 2.0 W/cm² in continuous mode and 3.0 W/cm² in pulse mode.

It is not recommended to increase in steps two parameters at the same time - if you extend the application area, do not increase intensity, and vice versa.

1.8 FREQUENCY OF TREATMENTS

Recommended frequency of treatments for acute conditions is 5 times a week and for chronic conditions, 3 times a week. It is advisable to change frequency during one set of treatments (for example, 3 times a day, and then 3 times every other day).

The total number of treatments varies according to the individual condition. It can range from a single application before myoskeletal surgery up to 9 applications over a 3-week period for chronic conditions.

1.9 COMBINED THERAPY

Combined therapy is the simultaneous application of two or more kinds of energy. (Common diadynamic currents with simultaneous application of galvanic and pulse low frequency currents can also be considered combined therapy.) Typical combinations are:

- ultrasound + low-frequency currents
- ultrasound + amplitude-modulated mid-frequency currents
- ultrasound + TENS

For combined therapy, we recommend using the CV mode of electrotherapy.

1.9.1 ULTRASOUND + LOW-FREQUENCY CURRENTS

This combination can be applied by using one BTL Combi unit (for example, BTL-4810S Topline Combi) or by the interconnection of two single units (BTL-4000 Topline Sono and BTL-4000 Topline Puls). The current with adjustable intensity is brought to the covering metallic plate of the emitter head. The other electrode (most usually a plate electrode) is placed contralaterally, so that both the ultrasound field and low-frequency currents are brought to the treatment area.
• **Mechanism of Action**

When using appropriate low-frequency current (100 – 200 Hz), the myorelaxation effect of the ultrasound is cumulatively increased. This form of combined therapy (when using ultrasound of frequency 3 MHz) is used especially for treating localized muscle hypertonia, uncoordinated muscle fibres (unable to spontaneously relax), and trigger points in superficially located muscles.

A certain disadvantage of this combination is the galvanic (corrosion) effect. This applies especially to diadynamic currents and to a certain extent also to monophasic pulse currents. This effect is more evident on the emitter head than on the skin. However, when using higher intensities, it is important to be aware of this effect and ensure the patient’s safety.

• **Intensity**

Set the intensity of ultrasound separately - 0.5 - 0.7 W/cm² for continuous ultrasound and up to 1.0 W/cm² for pulsed ultrasound with Duty factor 1:2 or more. The intensity of the low-frequency component is set according to the aim.

When locating trigger points, set the intensity at the threshold sensitive level outside the area of expected reflex changes and then approach this area continuously. In the area of the hyperalgesic zone, the pre-set intensity is above threshold sensitive (to just below the pain threshold), and the patient reports subjective intensification of feelings; may even feel pain. In the area of trigger points in a muscle, the intensity changes to the threshold motor level and up to above the threshold motor level. The pre-set intensity causes a motor response. (The patient often does not notice it.) If the pain threshold is not reached, use the same intensity for the therapy; otherwise, adjust the intensity of the low-frequency component according to the type of current and the desired effect. (The intensity of the therapy should never be above the pain threshold!)

The size of the indifferent plate electrode should be at least 2x (but in practice, even up to 10x) larger, so that sensitive feelings are located only on the edge of the ultrasound application.

1.9.2 **ULTRASOUND + AMPLITUDE MODULATED MID-FREQUENCY CURRENTS (2-POLE INTERFERENCE)**

Unlike the previous combination, mid-frequency currents have no galvanic effects. In addition, they are better tolerated (at the given intensity of subjective feelings, higher current intensity can be applied), the effect is deeper, and hence they can influence reflex changes in deep muscles.

It is obvious that for application to deeper areas, lower ultrasound frequency (1 MHz) is chosen. A combination of modern BTL units enables you to select ultrasound parameters or amplitude modulation. Hence, these currents can be used for the treatment of superficial (3 MHz frequency) as well as deep (1 MHz frequency) muscles.

• **Mechanism of Action**

It is the same as for the combination of ultrasound + low-frequency current (see the previous paragraph). The optimal myorelaxation is attained with a frequency of 150-180 Hz (unlike diadynamic currents, the myorelaxation effect is not lowered by the stimulant frequency of the 50 Hz - MF component of DD currents).

If you want to enhance myorelaxation of deeply located muscles by simultaneous increase in temperature, use continuous ultrasound. On the contrary, if increase in temperature is contraindicated, use pulsed ultrasound with a higher Duty factor (For Duty factor, see the chapter **Modes of Operation**).
• **Intensity**

The initial intensity for continuous ultrasound (Duty factor 1:1) is 0.4 - 0.6 W/cm², the initial intensity for pulsed ultrasound with Duty factor 1:2 - 1:4 is 0.5 - 1.0 W/cm², the initial intensity for pulsed ultrasound with Duty factor 1:8 - 1:16 is 1.0 - 1.3 W/cm².

The AMP intensity is chosen according to the desired effect. Mainly the analgesic effect of amplitude modulation about 100 Hz is reached by above threshold sensitive intensity, optimal myorelaxation effect of 150 – 180 Hz modulation is reached with intensity at the threshold motor level. The patient must not feel pain or a burning sensation during application even in the hyperalgesic zone or trigger point. If pain occurs, lower the ultrasound intensity immediately (record it on the patient’s card and send the patient to his/her physician for an immediate check-up). In order to eliminate the burning sensation in the trigger point area, lower the intensity of the mid-frequency component or add the contact medium (It is not necessary to inform the patient’s physician).

1.9.3 **ULTRASOUND + TENS (TRANSCUTANEOUS ELECTRICAL NERVE STIMULATION)**

In the above-mentioned combinations, the emphasis is put on myorelaxation, therapy of hyperalgesic zones and trigger points. In the combination of Ultrasound + TENS, the expected effect is generally analgesic, i.e. it is used not only for treatment of the above-mentioned reflex change areas, but also for other kinds of pain. It is important to keep in mind the mechanisms of ultrasound treatment because not every type of pain can be treated by ultrasound. This combination is especially suitable for myalgia (post-stress, reflexive, etc.) and post-traumatic pain (after peracute condition; 24-36 hours after onset), during lege artis general therapy (relaxation, etc).

The preferable TENS pulses are asymmetric, biphasic programs beginning with the numbers 12, 15, 18 (The stimulant effect of the ultrasound head/cathode is preserved and at the same time, the galvanic effect is eliminated). The continuous (or possibly random) frequency of 100 Hz is used.

The effect of this combination is better than the effect of the same treatments applied subsequently. It is necessary to be aware of all ultrasound contraindications (see the chapter **Specific Contraindication for Ultrasound Therapy**).

• **Mechanism of Action**

Reduction in the perception of pain positively affects the trophics of the injured area. The simultaneous application of ultrasound with consequent increase in capillary permeability, improved extravasal liquid absorption and thus reduction in pressure, increased vasoactive amines secretion, and direct effect on the precapillary sphincters are the best prevention against algodystrophic syndrome. Please note that beginning with the earliest clinical symptoms of algodystrophy, the application of any physical therapy to the porotic area is contraindicated, and only the segmental technique can be used (for example, ultrasound + electrotherapy applied paravertebrally, homolaterally on the segmental level, innervating the porotic area).

• **Intensity**

For the ultrasound intensity, please see the previous section. Intensity of TENS is above threshold sensitive outside the area of reflex changes and above the threshold motor level in the area of reflex changes.
1.10 SPECIFIC CONTRAINDICATIONS FOR ULTRASOUND THERAPY

1.10.1 ABSOLUTE

➢ Epiphyses Of Growing Bones

There is the risk of irreversible damage of the growth zone, deformity and permanent impairment. Since older types of ultrasound units employed low intensity (the displayed data did not correspond with the actual energy applied), damage to the growth zones was exceptional and many physicians did not take this contraindication too seriously. However, modern, effective ultrasounds can cause damage to the growth zone, especially due to insufficient movement of the emitter head. Responsibility lies with the physician who prescribed the therapy as well as with the therapist who performed the contraindicated procedure.

➢ Gonads

Small doses cause temporary and high doses permanent spermio/oogenesis impairment.

➢ Pregnancy

➢ Eyes

The use of special ophthalmic ultrasound units is restricted to specialised centres. In physiotherapy, it is contraindicated.

➢ Post-Laminectomy Conditions

After laminectomy, the spinal cord is not completely covered by the osseous cover, and ultrasound application can cause temporary or even permanent paraparesis of the lower extremities (if the axons of the spinal cord paths are disorganised). This is some of the most severe damage in physiotherapy practice and the link between the physiotherapy application and the condition is obvious.

➢ Bleeding With Recent Onset (Any Location)

Ultrasound waves propagate far via body fluids (though with significant energy loss). This can cause, for example, the re-occurrence of stubborn epistaxis (e.g., when the ultrasound is applied to a knee).

1.10.2 RELATIVE

➢ Brain, Parenchymatous Organs, Heart

There is no reason to apply ultrasound to these organs (The brain is covered by the cranium). In spite of this, there has been reported the case of a patient who irradiated his sinuses while he applied ultrasound to the knee and after a few hours died of massive bleeding into his forehead lobe (the calcified atheroma plate might have been disturbed). The patient must not apply ultrasound to himself/herself under any circumstances, not even to areas easy to reach!

➢ Peripheral Nerves (Situatned On The Bone, Just Below The Surface)

It mainly concerns n.ulnaris in the elbow area, volar areas of the wrist, inguines, areas under external and internal ankles, etc.

In all of the above-mentioned areas, constructive interference (stationary waves) results in local increase in intensity (peaks of intensity) and thus prolongation of impulse transmission and irreversible destruction of nerve fibres without damaging muscular fibres. Myeline covers of damaged nerves are usually preserved.
- **Bony Projections Located Under The Skin** - Spinous Projections Of Spondyles, Ankles, Condyles, Epicondyles
  
  This contraindication is very often ignored. Ultrasound can be used for the treatment of epicondylitis, but this is true only in case of applying ultrasound to a specific muscular group (extensoric in lateral epicondylitis, flexoric in ulnar epicondylitis). Direct ultrasound application to a painful attachment often causes increased pain, but it more often leads to the process of chronification (the same effect as in the case of repeated local application of corticosteroids).

  Similarly, direct ultrasound application to the spine often results in stubborn periosteum pain around spinal projections.

- **Emphysema, bronchiectasis (ultrasound application to the chest area)**

- **Menstruation**
  
  Ultrasound application is absolutely contraindicated in the lower abdomen area. When applying ultrasound to other areas, it is advisable to tell the patient that menstruation might increase (more common is acceleration of menstruation than metrorrhagia). If the case history shows that the ultrasound application might adversely affect menstruation, the therapist should refuse to carry out the treatment and suggest that the patient consult it with her physician. (It is not enough to postpone the ultrasound application after the menstruating period when ultrasound indications may not exist anymore.)

- **Blood circulation insufficiency**
- **Pacemaker**
- **Cardiovascular diseases**
- **Cochlear implants**
- **Metal implants**
- **Tumours**
- **Acute inflammations**
- **Vascular and blood diseases (haemophilia)**
- **Generally poor state of health**
- **Endocrine glands**
- **TB**
1.11 THERAPY PRESCRIPTION

- Name, surname, ID number, insurance company
- Diagnosis (name and number), stage
- Name of treatment, type of device, if possible (BTL-4000 Topline Sono, BTL-4000 Topline Puls + BTL-4000 Topline Sono)
- Ultrasound parameters:
  - frequency
  - duty factor
  - pulse frequency
  - intensity
  - emitter head size
  - way of application
- In case of combined therapy - parameters of electrotherapy:
  - characteristics of current
  - intensity
  - position and size of indifferent electrode
- Length of application, step
- Frequency of treatments, changes during the set of treatments
- Total number of treatments
- Date of check-up at the physician's
- Date of prescription, physician's name, signature, stamp

1.12 INSTRUCTIONS FOR THE PATIENT

Treatment is usually asymptomatic, a feeling of moderate warmth can occur. A burning sensation during therapy indicates insufficient contact medium (it is necessary to add more), pain indicates overdose (it is necessary to interrupt therapy and consult the prescribed treatment with the physician; treatment with lower intensity usually continues after a one-day break) or insufficient movement of the emitter head.

After the first treatment there may occur temporary worsening. Marked subjective and objective improvement should occur not later than after the third application. If there is no improvement (in acute and subacute states), ultrasound therapy should not be continued. Telling the patient that improvement will occur after a few months is not professional and it shows that the physician prescribed the treatment for the deferring effect (waits for the self-healing processes of the body).

If ultrasound is used for treating chronic conditions (which is quite rare) or for change in the mechanical features of treated tissues (e.g., in case of Dupuytren's contracture or plantar aponeurosis related to calcar calcanei), significant improvement cannot be expected after the third application. (If ultrasound application is not followed immediately by manual therapy, improvement cannot be expected even after the tenth application).
1.13 TREATMENT PROCEDURE

1. Inform the patient about the procedure.
2. **Never let patients apply ultrasound to themselves! It is strictly forbidden!**
3. Ask the patient about contraindications that may be temporarily present – epistaxis, menstruation or pregnancy.
4. Place the patient in the proper position for the therapy and apply the contact medium.
5. Set the prescribed intensity, time, place the emitter head and switch on the ultrasound.
6. Move the emitter head continuously according to the prescribed form of application.
7. Ask the patient repeatedly about his/her feelings. In case of a burning sensation, add more contact medium, and if pain occurs, lower the intensity or interrupt the treatment and inform the physician.
8. After the prescribed time has elapsed, lower the intensity and switch the unit off. (Most modern ultrasound units do it automatically).
9. Wipe the contact medium off the patient and dry the skin.
2 SETUP OF ULTRASOUND GENERATOR

2.1 SETUP AND CONTROL OF ULTRASOUND THERAPY

On the therapy parameters screen it is possible to select ultrasound or ultrasound sequences.

2.1.1 ULTRASOUND

- Carrier Frequency
  Select 1 MHz or 3 MHz. The 1 MHz frequency is used for deeper located tissues.

- Duty Factor
  Duty factor is the percentage ratio of the pulse length to the period length (the value in the brackets states the ratio between the pulse length and the period length). If the duty factor is set to 100% (1:1) the ultrasound works in so-called continuous mode. In the pulse mode, it is possible to select the duty factor from the following values: 6.25% (1:16), 12.5% (1:8), 25% (1:4) and 50% (1:2). For a description of finer setting of the duty factor and a more detailed description of the output signal course, see the section Course of signal.
  The thermic effects of ultrasound differ according to the duty factor value. Continuous ultrasound (duty factor 100%, i.e. 1:1) causes deep warming of tissues, while pulsed ultrasound (50% (1:2) and 25% (1:4)) has only slight thermic effects, and for the values 12.5% (1:8) and 6.25% (1:16) the thermic effect is totally zero.

- Intensity
  Intensity can be set within a range from 0.1W/cm² to 2W/cm² for the continuous mode, and from 0.1W/cm² to 3W/cm² for the pulse mode.
  If the continuous mode is set and then there is set an intensity higher than 2W/cm², the equipment sets duty factor to 50%. Similarly, if there is set intensity from 2.1W/cm² to 3W/cm² and then the duty factor is set to 100%, the equipment limits the intensity to 2W/cm².
• **Duration**

The duration of therapy can be set within a range from 00:01 to 30:00 [min:sec].

• **Pulse Frequency**

Pulse frequency can be selected from the following values: continuous, 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, and 150 Hz.

• **With Electro**

If your unit does not contain any generator of electrotherapy, the therapy parameters manual settings screen displays the button with electro. Pressing this button enables you to connect one pole of electrotherapy from the external connector to the case of the ultrasound head and perform combined therapy. For information about the configuration of output connectors and interconnection of units, see the BTL-4000 Topline Series User’s Manual. If your device contains an electrotherapy generator, this button is not displayed and one pole of electrotherapy is automatically connected to the case of the head after starting the combined therapy.

• **Course of Signal**

The course of signal button contains a picture informing about the shape of the modulation signal. For detailed information about the signal and for the option of fine setting the duty factor, press the course of signal button.

**Pulse [ms], period [ms]** — information about the length of the pulse and the period calculated for the set pulse frequency and duty factor.

**Pulse freq. [Hz]** — information about the set pulse frequency.

**Duty factor [Hz]** — via the select button it is possible to set a value from 6 to 100%. By pressing the duty factor button, you can select one of the preset values. See the section **Duty Factor**.

• **Physiological Effects**

The symbols of effects used in the equipment have the following meaning:

A  - analgesic
E  - antiedematous
F  - antiphlogistic
H  - hyperaemic
R  - myorelaxation

### 2.1.2 ULTRASOUND SEQUENCE

This function enables you to run sequences selected by name or number. It is also possible to create new sequences or edit existing ones.

Since sequence is a series of individual therapies one after the other, it is obvious that the parameters of individual sections must be set when creating the programs.

For details, see the chapter **USER SEQUENCES** in the BTL-4000 Topline Series User’s Manual.
2.2 SPECIFIC SETTINGS

The heads delivered with the BTL-4000 Topline devices are interchangeable. This means that the head can be connected to any device of the BTL-4000 Topline series which contains an ultrasound generator and it is not necessary to state the number of the equipment for which the head is designed.

For optimum detection of the contact between the head and the patient's skin, it is necessary to perform calibration of the ultrasound head, especially for extra purchased heads – see the chapter Ultrasound Head Calibration.

The ultrasound heads supplied with the equipment were calibrated during manufacture.

2.2.1 ULTRASOUND HEAD CALIBRATION

When calibrating the ultrasound heads, the equipment measures their properties so that optimum evaluation of the contact with the patient's skin is ensured.

- **Starting Calibration**

Put the ultrasound head in water and press the start button. The pilot lights of the head which is selected for calibration blink before the start of calibration and shine during calibration.

- **Finishing Calibration**

Take the ultrasound head out of the water, dry the metal part and press the finish button.

- **Saving Calibration Values**

To save the values measured during calibration of the head, press the save button.

To quit the head calibration function, press esc.

If you decide not to save the measured data (e.g. if the head was not immersed in water after the start of calibration), press the esc button to refuse saving.

After the end of calibration, start therapy for 1 MHz and 3 MHz (e.g. with intensity 1 W/cm²) and put the head in water and take it out several times, to check the correctness of contact detection.

**ATTENTION!**

If the prescribed procedure of ultrasound head calibration is not followed, there may occur damage to the ultrasound generator and the ultrasound head after the start of therapy.
3 TECHNICAL PARAMETERS

3.1 PARAMETERS OF ULTRASOUND GENERATOR

<table>
<thead>
<tr>
<th>Effective intensity</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Continuous operation</td>
<td>0.1 to 2W/cm² ± 20% for output intensity higher than 0.2W/cm²</td>
</tr>
<tr>
<td>Pulse operation</td>
<td>0.1 to 3W/cm² ± 20% for output intensity higher than 0.2W/cm²</td>
</tr>
<tr>
<td>Working frequency</td>
<td>1 MHz ± 5% and 3.2 MHz ± 5%</td>
</tr>
<tr>
<td>Modulation frequency</td>
<td>10 to 150 Hz ± 5%</td>
</tr>
<tr>
<td>Duty factor</td>
<td>6 to 100% ± 5% of the set value</td>
</tr>
<tr>
<td>Duty factor – defaults</td>
<td>6.25% (1:16), 12.5% (1:8), 25% (1:4), 50% (1:2), 100% (1:1) ± 5% of the set value</td>
</tr>
<tr>
<td>Time of therapy</td>
<td>0 to 30 minutes ± 2% of the set value</td>
</tr>
<tr>
<td>Maximum output power</td>
<td>12W</td>
</tr>
</tbody>
</table>

Table of pulse parameters

<table>
<thead>
<tr>
<th>Duty factor</th>
<th>Frequency 10 Hz period 100 ms</th>
<th>Frequency 50 Hz period 20 ms</th>
<th>Frequency 100Hz period 10 ms</th>
<th>Frequency 150 Hz period 6.67 ms</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Pulse length</th>
<th>Pause length</th>
<th>Pulse length</th>
<th>Pause length</th>
<th>Pulse length</th>
<th>Pause length</th>
<th>Pulse length</th>
<th>Pause length</th>
</tr>
</thead>
<tbody>
<tr>
<td>50%</td>
<td>50 ms</td>
<td>10 ms</td>
<td>5 ms</td>
<td>3.33 ms</td>
<td>3.33 ms</td>
<td></td>
<td></td>
</tr>
<tr>
<td>25%</td>
<td>25 ms</td>
<td>75 ms</td>
<td>5 ms</td>
<td>7.5 ms</td>
<td>1.67 ms</td>
<td>5 ms</td>
<td></td>
</tr>
<tr>
<td>10%</td>
<td>10 ms</td>
<td>90 ms</td>
<td>2 ms</td>
<td>9 ms</td>
<td>0.67 ms</td>
<td>6 ms</td>
<td></td>
</tr>
<tr>
<td>6%</td>
<td>6 ms</td>
<td>94 ms</td>
<td>1.2 ms</td>
<td>9.4 ms</td>
<td>0.40 ms</td>
<td>6.27 ms</td>
<td></td>
</tr>
</tbody>
</table>

3.2 STEP IN SETTING THE ADJUSTABLE VALUES

<table>
<thead>
<tr>
<th>Intensity</th>
<th>0.1 W/cm²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Modulation frequency</td>
<td>10 Hz</td>
</tr>
<tr>
<td>Duty factor</td>
<td>1%</td>
</tr>
<tr>
<td>Time of therapy</td>
<td>1 s</td>
</tr>
</tbody>
</table>
### 3.3 Parameters of Ultrasound Heads

<table>
<thead>
<tr>
<th>BTL-237-1-13 – small head</th>
<th>BTL-237-4-13 – big head</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Effective radiation area (ERA)</strong></td>
<td><strong>Effective radiation area (ERA)</strong></td>
</tr>
<tr>
<td>ERA (EN 61689)</td>
<td>0.7 cm² ± 20%</td>
</tr>
<tr>
<td>ERA (21 CFR 1050)</td>
<td>0.9 cm² ± 20%</td>
</tr>
<tr>
<td>Maximum effective intensity</td>
<td>3 W/cm² ± 20%</td>
</tr>
<tr>
<td>Maximum effective acoustic power</td>
<td>2.1 W ± 20%</td>
</tr>
<tr>
<td>Radiation frequency</td>
<td>1 MHz and 3.2 MHz ± 5%</td>
</tr>
<tr>
<td>Type of beam</td>
<td>collimated</td>
</tr>
<tr>
<td>Beam non-uniformity ratio (BNR)</td>
<td>&lt; 8</td>
</tr>
<tr>
<td>Covering grade according to EN 60 529</td>
<td>IP 67</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Effective radiation area (ERA)</strong></th>
<th><strong>Effective radiation area (ERA)</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>ERA (EN 61689)</td>
<td>3.2 cm² ± 20%</td>
</tr>
<tr>
<td>ERA (21 CFR 1050)</td>
<td>4.4 cm² ± 20%</td>
</tr>
<tr>
<td>Maximum effective intensity</td>
<td>3 W/cm² ± 20%</td>
</tr>
<tr>
<td>Maximum effective acoustic power</td>
<td>9.6 W ± 20%</td>
</tr>
<tr>
<td>Radiation frequency</td>
<td>1 MHz and 3.2 MHz ± 5%</td>
</tr>
<tr>
<td>Type of beam</td>
<td>collimated</td>
</tr>
<tr>
<td>Beam non-uniformity ratio (BNR)</td>
<td>&lt; 8</td>
</tr>
<tr>
<td>Covering grade according to EN 60 529</td>
<td>IP 67</td>
</tr>
</tbody>
</table>
4 ACCESSORIES

User's Manual
User's Guide
Head BTL-237-4-13 (large)
Head BTL-237-1-13 (small)
Ultrasound gel 300ml, optionally 1l
Touch-pen
Holder for touch-pen
Mains cable
Adapter
Built-in accumulator
Interconnection cable for combined therapy (optionally)
Carry bag (optionally)
Cart for BTL-4000 (optionally)