
Die Verhinderung lagerungsbedingter Schäden in der operativen Gynäkologie. Leitlinie der DGGG (S1-Level, AWMF-Registernummer 015/077, Februar 2015)

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Abstract

Purpose: Official guideline published and coordinated by the German Society of Gynecology and Obstetrics (DGGG). Positioning injuries after lengthy gynecological procedures are rare, but the associated complications can be potentially serious for patients. Moreover, such injuries often lead to claims of malpractice and negligence requiring detailed medical investigation. To date, there are no binding evidence-based recommendations for the prevention of such injuries.

Methods: This S1-guideline is the work of an interdisciplinary group of experts from a range of different professions who were commissioned by DGGG to carry out a systematic literature search of positioning injuries. Members of the participating scientific societies develop a consensus in an informal procedure. Afterwards the directorate of the scientific society approves the consensus.

The recommendations cover:
- responsibility, information provided to the patient, documentation
- basic principles to prevent positioning injuries
- prevention of specific types of positioning injuries such as injuries
  - of the upper extremities
  - of the lower extremities
- injuries caused by high-frequency surgery
- pressure ulcers caused by positioning
- prevention of compartment syndrome
- clinical monitoring and the diagnosis of positioning injuries

Zusammenfassung


Die Empfehlungen berücksichtigen:
- Zuständigkeiten, Aufklärung Dokumentation
- Grundsächliche Aspekte zur Prävention lagerungsbedingter Schäden
- Spezifische Aspekte zur Prävention lagerungsbedingter Schäden
  - der oberen Extremität
  - der unteren Extremität
- Schäden durch Hochfrequenzchirurgie
- Lagerungsbedingte Dekubitalulzera
- Aspekte zur Prävention des Kompartment syndroms
- Klinische Kontrollen und Diagnostik lagerungsbedingter Schäden

I. Guideline Information

DGGG Guidelines Program
Information on this topic is provided at the end of the guideline.

Citation format

Fleisch M et al. The Prevention of... Geburtsh Frauenheilk 2015; 75: 792–807
Guideline documents

The editorially complete, long version and a PDF slide set suitable for PowerPoint presentation of these guidelines as well as a summary of the conflicts of interest of all the authors can be found on the homepage of AWMF:

http://www.awmf.org/leitlinien/detail/ll/015-077.html

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See Table 1.

Abbreviations

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II Using this Guideline

Purpose and objectives

The reason for preparing the guideline was the increase in requests made to advisory bodies of German medical associations to review treatments in the context of long gynecological surgeries, particularly operations where patients were placed in the lithotomy position. As recommendations on the prevention and diagnosis of injuries are sometimes contradictory, a systematic review of the existing evidence on the prophylaxis, diagnosis and treatment of positioning injuries and a compilation of recommendations by a panel of experts was considered especially urgent.

Patient care

Outpatient and inpatient care.

Target audience

This guideline is addressed to the following groups of people:

▶ all physicians and professionals who treat gynecological patients intra- and postoperatively, in particular:
    ▶ gynecologists
    ▶ anesthesiologists
    ▶ nursing staff

Period of validity

The validity of this guideline was confirmed by the boards/responsible persons of the participating professional associations/working groups/organizations/societies as well as by the board of the DGGG and the DGGG Guideline Commission in February 2015 and thereby approved in its entirety. This guideline is valid from February 17, 2015 to February 28, 2018. The period of validity has been estimated based on the guideline’s contents. The guideline can be updated earlier if necessary; likewise, the guideline’s period of validity can be extended if it continues to mirror the current state of knowledge.
The methodology for the compilation of this guideline is prescribed by the classification assigned to the guideline. The AWMF Guidance Manual and Rules for Guideline Development (Version 1.0) sets out the rules for classifying guidelines. Guidelines are differentiated into lowest (S1), moderate (S2) and highest (S3) class. The lowest class of guideline is defined as consisting of a set of recommendations for action compiled by a representative group of experts from medical societies. In 2004 the S2 class is divided into two subclasses: s2e (evidence-based) and S2k (consensus-based). The highest class (S3) combines both approaches. This guideline is classified as: S1

In the planning stage the members of the guideline development group compiled a list of topics considered relevant for the “Prevention of Positioning Injuries during Gynecologic Operations”. A systematic literature search based on the list of topics/questions was done in PubMed using the following search terms (last updated on 31.01.2014): intraoperative positioning (mesh-term, n = 1582 hits), positioning damage (mesh-term, n = 672), positioning injury (mesh-term, n = 3120), intraoperative malpositioning (mesh term, n = 72), intraoperative posture (mesh term n = 1092), pressure ulcer (mesh term) AND prevention & control (subheading, n = 40) AND operation procedures (n = 47) OR peri-operative prevention AND devices (n = 82), compartment syndrome (mesh term) AND postoperative (subheading) (n = 1014).

After eliminating duplicates, publications in English or German were examined with regard to their relevance for the topic. Randomized studies, case control and observational studies, case reports, reviews, meta-analyses and Cochrane reviews were included. In principle, publications were only included if they referred to types of positioning or injuries which had a particular relevance in gynecological and obstetric surgery. Relevant recommendations and guidelines on the topic issued by the following societies were also considered:

- The agreement on cooperation in gynecologic and obstetric surgery issued by the German Society of Anesthesiology and Surgical Intensive Care (DGAi) and by the Professional Association of German Anesthesiologists (Berufsverband Deutscher Anästhesisten [BDAi]) together with the German Society for Gynecology and Obstetrics (DGfG) and the (German) Professional Organization of Gynecologists [1]
- “Recommended Practices for Positioning the Patient in the Perioperative Practice Setting” issued by the Association of Perioperative Registered Nurses (AORN): [2]
- "AST Recommended Standards of Practice for Surgical Positioning" published by the Association of Surgical Technologists [4] (AST)
- The practice guidelines (Expertenstandard) issued by the German Network for Quality Development in Nursing (Deutsches Netzwerk für Qualitätsentwicklung in der Pflege [DNQfP]) [6] on preventing pressure ulcers (Dekubitusprävention)

The current guideline and its synthesized recommendations for the prevention of positioning injuries were adopted by all 14 members of the Commission following an informal consensus process.

### 2 Preamble

Preoperative positioning of the patient and appropriate intraoperative positioning during gynecologic operations is an interdisciplinary task which requires the cooperation of professionals across a range of disciplines, all of whom have a legal obligation of due care. The aim is to ensure patient safety and prevent position-related injuries. It is also necessary to balance the surgeon’s concerns regarding the best view of the surgical site through positioning and maneuvering the patient against the concerns of the anesthesiologist about maintaining the best and safest means of access to the patient. Moreover, the patient’s dignity should be protected throughout the procedure.

Incorrect positioning can damage the patient’s health; such damage can be transient but some injuries are permanent and can result in long-term functional restrictions, secondary morbidity, and even death.

Optimal patient positioning should prevent pressure injuries (pressure ulcers), skin irritation, burns, nerve damage, circulatory problems and hypothermia. Positioning injuries can affect the skin and soft tissues, joints, ligaments and bones as well as the eyes, nerves and vessels.

### 3 Epidemiology

Because of muscle relaxation, patients under anesthesia lack protective reflexes and muscle tone. They are at increased risk of injury, in particular injuries such as joint luxation, plexus injuries and pressure ulcers. Moreover, around 30% of all patients complain of back pain postoperatively, irrespective of the anesthetic technique used for the operation [7], with up to 37% of patients complaining of back pain after undergoing surgery in the lithotomy position [8].

The overall incidence of pressure injuries reported in the literature is 5% [9]. The most common patient positions used in gynecologic surgery are the supine position, various modifications of the lithotomy position [10] with or without Trendelenburg positioning, and the upright seated position (“sit-up” or “beach chair” position) for breast surgery [11]. Nowadays the lithotomy position is usually modified; nevertheless it is associated with a particular risk of positioning injury. The number and duration of operations carried out with the patient in an extreme Trendelenburg position to improve exposure of the lesser pelvis in laparoscopic or robotic-assisted procedures has increased and with it the risk of positioning injuries [12, 13]. The Trendelenburg position is used in these procedures to shift the abdominal viscera from the pelvis cranially to improve exposure.

Between 2008 and 2012 the advisory bodies of the regional medical associations in Germany reviewed 63 procedures in connection with charges of positioning-related injury in gynecology and obstetrics (source: German Medical Association). This corresponds to 1.8% of all accusations of medical error in gynecology in Germany. In 31.7% of cases the advisory bodies confirmed medical error (overall percentage for gynecology: 32%). This percentage is higher than that of comparable procedures where there is a suspicion of positioning-related injuries during surgery in other medical specialties (22.5%). The reviewed positioning injuries occurred almost exclusively after operations performed with the patient in a (modified) lithotomy position.

Positioning injuries during surgery are the cause of around 4% of all medical errors confirmed by the advisory bodies of German
4 Basic Forensic Aspects of Positioning

4.1 Assignment of tasks and responsibilities

The responsibility for positioning the patient preoperatively, intraoperatively and postoperatively is defined by the agreements of the professional societies (with the DGAI and the BDA) [1]. Positioning in the operating room (OR) is divided into four stages with different persons considered responsible for correct positioning:

1. Preoperative stage: The anesthesiologist is responsible for positioning the patient to administer the anesthesia and for monitoring the patient until the patient is properly positioned for surgery.

2. Intraoperative positioning: The surgeon is responsible. The anesthesiologist must alert the surgeon if there are visible errors or concerns. The anesthesiologist is responsible for positioning the extremities which are required to monitor the level of anesthesia and to administer anesthetics and infusions. The anesthesiologist is responsible for taking specific and appropriate precautions when positioning the patient to monitor and maintain the patient’s vital functions.

3. Deliberate intraoperative change of positioning: The decision and responsibility rests with the surgeon (although the change may be initiated by the anesthesiologist). The surgeon must be alerted to unintended changes in positioning, and the surgeon must then take the decision on how to proceed and is responsible for this decision.

4. The anesthesiologist is responsible for positioning including repositioning of the patient after surgery up until the patient has fully recovered from the anesthesia, unless special circumstances require the involvement of the surgeon when repositioning the patient.

In all 4 stages, a qualified surgical nurse or surgical anesthesia assistant can be directed to do the actual positioning of the patient. However, at every stage, the responsibility still lies with the physician, even if the person doing the positioning is a qualified specialist for patient positioning. The qualified positioning specialist is responsible for positioning the patient properly but cannot be held liable for the decision about the position he has been directed to place the patient. A lack of general instructions regarding the type of position and a lack of monitoring of the patient’s position is considered an organizational mistake (ruling by the German Federal Court of Justice on 24.01.1984 – VI ZR 203/82 – loc.cit.)[14].

The relevant agreements concluded between the respective professional associations always emphasize that a separation of responsibilities with respect to preoperative, intraoperative and postoperative positioning of patients must only be understood in the spirit of carrying out shared tasks on behalf of the patient [1]. But in the event of disputes regarding liability, the precise assignment of the area of responsibility will take priority, meaning that every medical specialty and each professional group must take steps to ensure that the assigned areas of responsibility are shown clearly in the documentation. The decision about the type of position is taken based on what is required for the surgical procedure after taking the anesthesia risk into consideration [1].

If the responsible anesthesiologist has concerns regarding a particular type of position because it limits monitoring or may affect the maintenance of vital functions or may be associated with a risk of positioning injury, then he must alert the surgeon to his concerns [1,15].

In the rare case that the surgeon and the anesthesiologist cannot agree on the type of positioning, then the so-called “casting vote” principle applies, where the surgeon has the decisive vote as the primary attending physician [15]. If, in such a case, the surgeon sticks to his assessment, then it must be assumed that he has done so after weighing up the competing interests [15]. In this case, the surgeon bears the professional and legal responsibility for ensuring that “surgical reasons justify the increased risk associated with his preferred positioning ...” [1].

4.2 Informing the patient

According to sec. 630d BGB (German Civil Code) the treating physician must obtain the patient’s consent prior to carrying out medical treatment. According to sec. 630d para. 2 BGB the effectiveness of this consent is contingent on the patient having been informed before giving consent. Sec. 630e para. 1 BGB requires the treating physician to inform the patient of all and any circumstances relevant for consent, including in particular the nature, extent, implementation, anticipated consequences and risks involved in the treatment, unless the patient has expressly waived her right to receiving such information. The risks of positioning include pressure injuries to soft tissues, nerve lesions and compartment syndrome.

The German courts have repeatedly considered the issue of positioning injury. According to a ruling of the Higher Regional Court (Oberlandesgericht, OLG) of Cologne, the assumption that “if the patient is technically positioned correctly on the operating table, the relevant rules for positioning are complied with, and patient’s position is monitored by the surgeon ... such measures are ... ‘entirely controllable’ is by no means automatic and does not apply to every type of positioning-related injury. In particular, pressure necrosis can often not be entirely avoided, even with the utmost care ...” (OLG Köln, ruling on February 25, 2013, Az. 5 U 152/12, MedR 2014, 399).

One of the insights of medical science is “that despite taking the utmost care when positioning a patient, positioning injuries can nevertheless occur” (OLG Koblenz, ruling on October 22, 2009, Az. 5 U 662/08, MedR 2010, 416). Even “optimal positioning ... does not always protect against developing pressure ulcers”, according to the Higher Regional Court of Hamm (OLG Hamm, ruling on May 20, 2011, Az. I-26 U 23/10).

Although the German Federal Court of Justice has stated that “the technically correct positioning of the patient on the operating table and compliance with the medical rules developed to protect patients from possible positioning injuries ...” are measures which belong to the “risk area of the hospital and of the medical staff ...” and which are “entirely controllable by the nursing staff and the responsible physicians”, this is not inconsistent with the physician’s obligation to inform the patient of the potential risk of positioning injuries. While technical methods for positioning the patient may be controllable, nevertheless, particularly in long operations, neither the positioning technique employed nor the monitoring of the patient’s position can completely prevent a change in the patient’s position, meaning that the positioning-related risk of neuropathies, pressure injuries, etc. cannot be completely excluded. This applies even more when patients have rare anomalies which were not detected despite complying with the current standards of care during the preliminary examination of the patient and when such anomalies facilitate pressure injuries.

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Important forensic aspects:
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still occur, despite evidence of monitoring and readjustment of
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When patient-specific risk factors are present or with certain
"Positioning is an interdisciplinary task which requires the co-
table and must be accessible. The staff entrusted with patient po-
tioning (nurses, surgeons, anesthesiologists for their area of
responsibility) must be trained in these standards.

Documentation of patient positions can be done with refer-
ce to the hospital’s recorded mandatory in-house positioning
standards. Deviations from these standards must be docu-
mented. If there are no mandatory in-house standards, a de-
tailed description of the patient’s position and any aids used
(gel mats, etc.) must be entered into the records and the surgi-
cal report.

Intraoperative monitoring by the surgeon to ensure the patient
is still correctly positioned does not have to be specially docu-
dented each time; however, it is expedient to record having
performed routine controls in the surgical report [15].

The patient’s position after an intraoperative change in posi-
tion (e.g. repositioning the patient from the classic lithotomy
position to a flat lithotomy position) is the responsibility of
the surgeon. It must be controlled and the repositioning must
be documented. The extent and type of control is not specified.

4.3 Documentation
The medical practitioner bears the burden of proof in terms of
showing that the patient was placed in a technically correct posi-
tion, with the medical practitioner having to clear himself of the
charge of malpractice in the event of positioning injury. This
means that providing evidence of having proceeded appropri-
ately and correctly is particularly important. Evidence is usually
provided by registering appropriate standards which take the
form of instructions on standard patient positions used in gyn-
ecology and obstetrics as well as specifically documenting the
measures taken for each patient, including identifying each per-
son who treated the patient. This documented information is
generally included in preprinted form on the surgical records,
which are usually collected and stored digitally and must be ini-
tiated by the person doing the positioning and the responsible
physician. It is not necessary to write a detailed report on posi-
tioning the patient during surgery (BGH ruling on January 24,
1984 – VI ZR 203/82 – ArztRecht 1984, 238). Instead, if the stan-
dards have been registered, all that is required is to report and
justify any individual deviations. When undertaking longer oper-
atations, it is recommended that areas likely to react sensitively to
longer patient positioning times are checked and the findings
documented; particularly if there is an intraoperative change of
the patient’s position, compliance with the respective standards
must be recorded in the surgical report. If positioning injuries
still occur, despite evidence of monitoring and readjustment of
the patient position, then this is an indication that these were ex-
ceptional circumstances which, in individual cases, would qualify
the assertion that the risk that occurred could have been entirely
manageable.

Important forensic aspects:

- Positioning is an interdisciplinary task which requires the co-
  operation of professionals across a range of disciplines.
- Individual responsibility for positioning is divided into differ-
  (surgeon), deliberate intraoperative change of position (sur-
  geon), postoperative p. (anesthesiologist).
- When patient-specific risk factors are present or with certain
  types of positioning for surgery which are considered to have
  an inherent risk for positioning injury (for example and pri-
  marily long operations performed with the patient in a lithot-
  omy position), patients should be informed by their physician
about risks of specific potential positioning injuries (e.g. com-
artment syndrome).
- The standards for typical patient positions used in gynecologic
interventions should be defined and recorded for every hospita-
l and must be accessible. The staff entrusted with patient po-
tioning (nurses, surgeons, anesthesiologists for their area of
responsibility) must be trained in these standards.

5 General and Specific Aspects of Correct Positioning
General positioning recommendations for all types of positioning
(“Good Practice Points”):
- It is recommended that surgical units develop positioning
standards, communicate them to the different medical special-
ties and to all professionals and staff, store the documented
standards and revise them at regular intervals to ensure the
contents are still up-to-date and take account of recent evi-
dence.
- Positioning materials in sufficient quantities and of a sufficient
quality must be available for the operation. The type and ex-
tent of materials purchased is determined by the patient pop-
ulation and the state of scientific knowledge. Particularly when
positioning obese patients it is important to ensure that the
operating tables have the appropriate weight specification to
support patients.
- Co-morbidities and pre-existing conditions which are relevant
for positioning must be determined preoperatively and must
be taken into account when positioning the patient (AORN).
These include endoprostheses and implanted devices, limited
joint mobility and anatomical anomalies (insofar as these are
known preoperatively).
- Positioning and repositioning, if done, must be carried out by
adequate numbers of personnel (AST, AORN) to ensure patient
safety and the ergonomic safety of staff.
- When the patient is transferred to the operating table, the pa-
tient must not be slid onto the table; the patient should be
transferred using appropriate transfer devices (back boards,
lifting) and with as little friction as possible to avoid injuries
to the skin from shear or friction (AST).
- It is important to ensure that the pad on which the patient lies
is dry and crease-free (minimize the number of sheets placed
under the patient).
- No parts of the body should hang over the edge of the operat-
ing table; the sacrum should not extend beyond the edge of the
operating table (AST, AORN).
Practice Points

General recommendations for the lithotomy position (positioning-related injury (Tables 2 and 3)): Provide intraoperative protection against hypothermia, with active warming of the patient using warming systems (WarmTouch, Bair Hugger) where necessary.

Pad the head to ensure that the cervical spine is in a neutral position and no pressure points develop at the back of the head.

If the arms are placed in an abducted position, abduction to approx. 60° should be done with the arms in a neutral position; any further abduction requires the arms to be in a supination position. The arms should be slightly flexed at the elbow and the forearm supported by an arm supporting device. The elbow joint should be placed on a pressure-minimizing padded support, the elbow should be at least at shoulder level (AST, AORN).

Both leg holders must be padded and at the same height (AST, AORN).

Depending on the patient’s physical constitution (obesity, joint mobility, etc.) the legs must be lifted into the leg holders and out of the leg holders at the end of surgery by sufficient numbers of personnel to prevent lumbosacral injury and hyperflexion of the hips (AST, AORN).

It is not known how long patients can remain in the lithotomy position without suffering positioning injury and the time probably depends on the physique and condition of the individual patient. The time spent in the lithotomy position should therefore be as short as required for the individual surgical intervention (AORN).

If the arms are placed alongside the body, it is important to watch out for the position of the fingers to prevent crush injury when moving the leg holders (AORN).

6 Prevention of Typical Positioning-related Injuries in Gynecology

Based on the available literature (observational studies) the following factors are associated with an increased incidence of positioning-related injury (Tables 2 and 3):

- BMI < 20 or > 30 [12]
- Diabetes mellitus [20, 21]
- Limited physical mobility (osteoarthritis, arthritis, ankylosing spondylitis, knee and/or hip endoprothesis, arthrodesis etc.)
- Age > 70 years [20]
- Malnutrition [22]
- Peripheral arterial occlusive disease
- Smoking and COPD [12, 22, 23]
- Anatomical abnormality (cervical rib, etc.) [24]
- Pre-existing neuropathies [12]

It is recommended that patient factors and circumstances which directly affect positioning in practice (e.g. endoprothesis, arthrodesis etc.) should be noted on the surgery plan and communicated to other staff involved early on.

6.1 Positioning-related neuropathies

The overall incidence of postoperative neuropathies (including lesions caused by surgery) is reported to be 0.6–1.2‰ [9]. However, in a study of 1210 patients who underwent major pelvic surgery the rate of postoperative neuropathies was 1.9% [27]. Neuropathies are generally caused by a combination of stretch, ischemia and pressure [28].

For practical purposes it is useful to classify the degree of severity of nerve damage into “neurapraxia”, “axonotmesis” and “neurotmesis”. This classification is important for the prognosis of nerve damage and for decisions concerning potentially necessary therapy. Neurapraxia and axonotmesis can occur in the context of positioning injury [29, 30]. Neurapraxia, which represents the lowest degree of severity, results in focal blocking of impulse conduction and is limited to the affected nerve segment. It can be caused by a brief interruption of blood supply due to external compression. Regeneration can take hours, days or weeks, in rare cases even months [29, 31]. Recovery after neurapraxia is complete and without sequelae. Axonotmesis is a traumatic injury with disruption of axons. The myelin sheath of Schwann cells also suffers injury but the connective tissue framework of the nerves (endoneurium, perineurium, epineurium) is preserved. After Wallerian degeneration of the axonal stump distal to the injury, the proximal axon stump forms a growth cone. The growth cone grows distally to the injury at a speed of 1 mm/day inside the preserved endoneurial sheath. If regeneration is successful, recovery of function is probable [29, 31]. However, the likelihood of com-
with respect to the shifting of patients intraoperatively and found the incidence of brachial plexus injury has not yet been demonstrated [45]. Whether the use of vacuum mattresses could reduce the in-"non-anesthetized patients, showed that use of a medical vacuum patient and simultaneously measured the pressure exerted on three different systems to prevent intraoperative sliding of the patient and the operating table [33,45]. The pressure exerted on the shoul-der appears to increase the risk of plexopathies [46]. Pressure increased [45]. The combination of arm abduction and shoulder extension together with rotation of the cervical spine [42]. Symp-toms of this plexus injury include motor and sensory deficits in the shoulder, the upper arm and forearm and the hands. Anatom-ical variants in the patient such as the existence of a cervical rib, the shoulder, the upper arm and forearm and the hands. Anatom-ical variants in the patient such as the existence of a cervical rib, an abnormal course of the plexus or deformity caused by fracture are all risks predisposing to brachial plexus injury [24,43]. The prognosis for full recovery of function after such injuries is gener-ally good, with a high probability that motor and sensory symp-toms will resolve over time [44], although recovery can take sev-eral months. Nevertheless, cases with permanent functional im-pairment have also been described in the literature [33].

In the steep Trendelenburg position the patient is head-down in a supine position with her feet 30° higher than her head. Shoul-der braces are often used to prevent the patient from sliding on the operating table [33,45]. The pressure exerted on the shoul-ders increases as the tilt angle in the Trendelenburg position is increased [45]. The combination of arm abduction and shoulder brace appears to increase the risk of plexopathies [46]. Pressure on the peripheral accessory nerve can additionally lead to trape-zius muscle paresis. One study, which prospectively evaluated three different systems to prevent intraoperative sliding of the patient and simultaneously measured the pressure exerted on the shoulder with varying degrees of head-down tilt angle in non-anesthetized patients, showed that use of a medical vacuum mattress system resulted in the least pressure on the shoulders [45]. Whether the use of vacuum mattresses could reduce the in-cidence of brachial plexus injury has not yet been demonstrated, even though many authors advocate the use of such systems. An-other study compared anti-skid foam mattresses with gel pads with respect to the shifting of patients intraoperatively and found no difference between systems [47].

Recommendations for prevention (Figs. 1 to 4):
- Minimize the degree of tilt and the time spent by the patient in a head-down position after taking the surgical aspects into consideration.
- Position the head using an appropriate device; avoid longer pe-riods of hyperextension or lateral flexion/rotation; limit arm abduc-tion to less than 90° (AORN, ASA) [28].
- Arm braces must be positioned in such way that dropping of the shoulder is prevented [28].
- Intraoperative sliding on the operating table should be avoided (AORN); combined with shoulder braces, the use of non-sliding operating table pads or mattresses decreases the pressure ap-plied to the plexus. This combination should be used in prefer-ence to shoulder braces alone.
- If shoulder braces are used, they must be padded and the point of contact should be at the level of the acromioclavicular joints (AORN).
- If shoulder braces are used, avoid or minimize additional ab-duction of the arms. Arm abduction must never exceed > 90° [48].

Ulnar neuropathy: Because of its largely unprotected course in the ulnar nerve sulcus, the ulnar nerve is at risk of pressure injury [49]. A prospective study of 1502 patients reported an incidence of 0.5% [50]. Analysis showed, however, that the risk population consisted primarily of men. This circumstance along with the fact that clinical symptoms only occurred between two and seven days after surgery suggests that factors other than inadequate pa-tient positioning may play a role.

Clinical symptoms of injury include paresthesia of the fourth and fifth finger and on the ulnar side of the hand. The full clinical pic-ture with involvement of ulnar nerve motor fibers results in the so-called ulnar claw [30]. Pressure injuries can occur through di-rect pressure in the area of the elbow created by incorrect posi-tioning of the arm, non-physiological pressure placed on the arm by the surgeon leaning on the patient, or pronation of the arm on the arm braces [30]. Typical findings on neurography consist of partial conduction block without accompanying conduction de-lay with and without axonal injury (depending on the duration and severity of the pressure injury) or loss of motor units on elec-tromyography (EMG). Depending on the severity (neurapraxia or axonotmesis), remission of paresis can take up to one year [51, 52].

6.1.2 Neuropathies of the lower extremity

According to a retrospective analysis of patients treated at the Mayo Clinic (Rochester/USA), persistent (≥ 6 months) motor neuropathies of the lower extremity occurred in around 1 of 3600 procedures in the lithotomy position [12]. In this study, every hour the patient remained in the lithotomy position increased the risk for neuropathy by a factor of 100. In 78% of cases the per-neal nerve was affected; the sciatic nerve was affected in 15% of cases and the femoral nerve in 7%. Sensory neuropathies oc-curred in 15: 1000 cases [12, 13]. Complete regeneration within the space of one year occurred in less than half (43%) of all cases [12]. On multivariate analysis, risk factors included BMI of 20 or less, a history of smoking within 30 days prior to the procedure, and prolonged duration of lithotomy of 4 hours or more.

Peroneal neuropathy: The common peroneal nerve, a division of the sciatic nerve, crosses the knee joint at the lateral aspect of the fibular neck [30] and divides into two branches. The superficial peroneal nerve provides sensory information but also innervates the peroneus longus and peroneus brevis muscles required for pronation of the foot. The deep peroneal nerve innervates the...
muscles required to lift the foot at the ankle and the arch of the foot and provides sensory information for the web space between the first and second toe. Because of the limited soft tissue cushioning at the fibular neck there is a risk of direct pressure injury. Pressure is often caused by unpadded contact with the leg holder. Alternatively, a combination of hip flexion and knee extension can lead to non-physiological traction of the sciatic nerve and the peroneal nerves [53]. Postoperative symptoms of common peroneal neuropathy include sensory deficits in the lateral lower leg and of the arch of the foot. Injury can result in motor deficits affecting the dorsal flexion of the foot, which may present clinically as drop foot. The differential diagnosis should include peroneal/sciatic injury and injury of a branch of the lumbosacral plexus which can occur in the lithotomy position [14]. Low BMI, smoking and prolonged duration of intervention increase the risk [12].

Recommendations for prevention (Figs. 5 to 7):

- Avoid direct pressure on the peroneal neck; pad potential pressure points.
- When using stirrups with straps, the leg should not be in contact with the rods of the support (AORN).

Sciatic neuropathy: Sciatic neuropathies have been described after the patient was in a lithotomy position and after cesarean section [54,55]. The lithotomy position can result in overextension of the peroneal aspect of the sciatic nerve [53,56,57]. Sciatic nerve lesions can lead to paresis below the knee. If knee flexion is...
preserved, the nerve is largely intact [58]. Injury can also lead to hypesthesia in the lateral aspect of the calf and across the entire area of the foot with the exception of the inner side of the foot. Recommendations for prevention (Figs. 5 to 7):

- Avoid overextension of the ischiocrural musculature; hip flexion should not exceed > 90° [3].

**Femoral neuropathy:** Several gynecologic case studies have reported on femoral neuropathies after procedures in the lithotomy position [59], some of them caused by self-retaining retractor systems [60]. In each case, hip abduction and extreme hip flexion with external rotation were cited as increasing the risk of injury [61–65]. Such positioning results in mechanical bending of the femoral nerve which is pressed against the inguinal ligament. In vaginal procedures with the patient in the lithotomy position, this mechanism may even be intensified by the surgical assistant leaning against the inner aspect of the thigh [28]. Clinical symptoms of femoral neuropathy include postoperative deficits in hip flexion and knee extension in combination with a reduction of the patellar reflex. The use of split leg tables, where the legs are in a supine position with abduction of both hips, is associated with femoral neuropathy. In robotic-assisted procedures where abduction was 25° the incidence for this complication was reported to be 1.7% [66]. Common symptoms are numbness of the legs and a tendency to fall when walking. Most sensory deficits disappear within five days [67]. In one case study, 94% of patients with motor symptoms experienced complete remission within 10 weeks; the remaining patients experienced remission within four months [68].

Recommendations for prevention (Figs. 6 to 8):

- Avoid hip flexion of > 90° in the lithotomy position; otherwise limit the time in this position.
- Avoid extreme abduction and external rotation of the hip.
- The surgical assistant must not lean against the inner aspect of the patient’s thigh (AORN).

**Obturator neuropathy:** The obturator nerve is a nerve of the lumbar plexus. There are few reports on obturator neuropathy after procedures in the lithotomy position [69]. Experiments have shown that abduction of the thigh between 30–45° in the hip joint results in significant traction of the obturator nerve which can be compensated for by hip flexion [70]. Obturator nerve injury caused by pressure of the fetus on the internal pelvic wall have been described after vaginal delivery [71]. This phenomenon must also be considered in the differential diagnosis of puta-
tive positioning-related obturator neuropathy after cesarean section.
Recommendations for prevention (Figs. 5 to 7):
- Abduction of the lower extremity of > 30° (lithotomy position or split-leg table) must be combined with flexion of the hip to prevent positioning-related neuropathy of the obturator nerve. The abduction angle should not exceed 45°.

6.2 Postoperative non positioning-related neuropathies
Occasionally (epidemiological data are lacking), peripheral neuropathies may develop after surgery where the anatomical pathology cannot be easily explained by the procedure. Such neuropathies are usually painful or start with the patient experiencing pain followed by paresis.
Such neuropathies correspond to the clinical picture known as neuralgic shoulder amyotrophy or plexus neuritis. Neither the losses in sensory perception, which are often minimal, nor the paresis can be explained by perioperative mechanical factors affecting the peripheral nerves such as traction, pressure, or sharp dissection. In many of these patients infection is considered the cause [72], although it remains entirely unclear how surgery is a causal factor for the infection. It is not possible to give evidence-based causal recommendations for treatment, although there are reports on the use of immunoglobulins and steroids. The neurological complaints generally improve over time, but complete remission is rare. It is clear that differentiating between these postoperative inflammatory neuropathies and the above-described neuropathies caused by mechanical factors is extremely important, both clinically and forensically.

6.3 Injuries caused by high-frequency surgery
High-frequency surgery is used in most gynecological procedures for cutting or coagulation. High-frequency alternating current flows from the active electrode through conductive tissue to a larger passive electrode (neutral electrode) which conducts the current back to the high-frequency unit again, thereby closing the electric circuit. Thermal energy is created at the point of contact between the active electrode and the tissue, which creates the desired cutting or coagulation effect. Technical or operating mistakes and surgery- or patient-related factors (e.g. uncontrolled loss of bodily fluids, amniotic fluid, etc.) can potentially endanger both patient and user. If the current density under the neutral electrode is too high at any point, this can lead to the unintentional release of thermal energy [14], which can spread unnoticed from the neutral electrode. Burns mainly occur with single-surface electrodes where it is not possible to monitor the quality of the contact between electrode and patient. Liquid bridges or a point of contact between the patient’s body and conductive material can result in leakage current. No high-frequency generator can measure and thereby avoid such leakage current. The Medizin-Produkte-Betreiberverordnung (German regulations governing the installation, operation and use of medical devices, MPBetreibV) stipulates that every user must be trained in the proper operation of and the risks associated with high-frequency units.
Prior to every application, the material used must be checked for defects.
Suspicious skin lesions which are noted postoperatively are not necessarily always associated with high-frequency current as they can also be caused by heat, pressure, time, chemicals and/or moisture (s. paragraph 6.4).
Recommendations for prevention:

- The patient must be placed on a dry, insulated support; wet sheets and underlays must be replaced by dry ones.
- The patient must not be in contact with electrically conductive surfaces.
- Avoid any puddles of disinfectants because of the risk they may be ignited by sparks (HF surgery).
- The full-sized neutral electrode should be placed near the surgical site preoperatively while maintaining sterility.
- The entire surface of the neutral electrode should be in contact with the patient’s skin; shave hairy areas if necessary.
- Ensure there are no traces of liquids between the skin and the electrode; do not use additional gel.
- Ensure that urine will be drained if the patient is scheduled for a lengthy surgical intervention (> 3 hours).
- The use of two HF units with two separate neutral electrodes in parallel is not recommended because of the potential that power output will briefly spike, creating a higher risk of burns for the patient.
- Remove all jewelry preoperatively; if an item cannot be removed, cover it with insulating tape and ensure that there is no contact with HF current and no high-frequency current is applied in the immediate vicinity of the area with the jewelry.
- There must be no item of jewelry in the area between the active and the neutral electrode.

6.4 Positioning-related pressure ulcers
Pressure ulcers are undesirable complications of surgical procedures which could, in principle, be avoided. They are the cause of additional suffering for affected patients (pain), increase the time spent in hospital, and involve additional treatment costs (for materials and personnel to treat the injury).

In 2009, the European and US Pressure Ulcer Advisory Panels published a guideline on pressure ulcer prevention which included a generally accepted definition of pressure ulcers:

“A pressure ulcer is a localized injury to the skin and/or underlying tissue usually over a bony prominence, as a result of pressure or pressure in combination with shear.” (European Pressure Ulcer Advisory Panel [EPUAP] and National Pressure Ulcer Advisory Panel [NPUAP], 2009).

Clear evidence of the causes of the skin injury is an essential part of the definite diagnosis (also to differentiate it from other skin injuries).

Pressure ulcers often develop in underlying tissue (just above the bony prominences in the musculature) while the overlying tissue layer initially remains intact. The injury may only become visible several days after it began to develop (i.e. during surgery).

6.4.1 Risk factors for the development of pressure ulcers
In addition to the causes of pressure ulcer described in the above definition (pressure and shear) other risk factors are also often discussed; however, their significance is not yet clear. More than 100 risk factors have been discussed in the literature [73]. Risk factors proposed as potentially causative in the perioperative period include: diabetes mellitus (OR = 2.15 [1.62–2.84]) [21], duration of anesthesia and total duration of hypotonia (< 50 mmHg diastolic blood pressure) [74], age > 71 years, dehydration, excessive skin moisture, nutritional deficiencies, sensory perception disorders, lung disease [22], central or peripheral nerve block (perioperative analgesia) [75], hypothermia [76], hypotonia, vascular disease, smoking, COPD [23], patient position during surgery (a lateral position is associated with a higher risk than a supine position, OR = 8.1) and duration of surgery (OR 3.7 for every doubling in the length of surgery) [25].

The methodological quality of the studies seeking to uncover (causal) connections between the risk factors listed above and the incidence of pressure ulcers is very heterogeneous and the overall level of evidence is rather weak. There are very few studies of patient populations in a gynecological surgery setting.

6.4.2 Perioperative prevention
Risk assessment: There is some disagreement as to whether a structured risk assessment approach is absolutely necessary for the effective prevention of pressure ulcers. The instruments employed in the assessment can yield false-positive or false-negative results, and there are certain situations (such as lengthy gynecological operations) in which every patient must be treated as a potential high-risk patient. Anesthesia-induced immobility and certain positions (sitting up, lateral position) increase the pressure in the tissue at risk.

A Cochrane review of recent studies on the use of pressure ulcer risk scales to reduce the incidence of pressure ulcers showed no benefits from the use of such scales [77].

Pressure-relieving devices: The choice of device depends on the functional deficits of the patient, i.e. the extent of pressure or shear exerted on the specific body part. Depending on the extent and duration of tissue pressure, pressure on affected areas must be relieved through positioning the patient on a soft mattress to prevent/minimize cell damage. The frequency of repositioning will depend on the support surface and the overlay (EPUAP/NPUAP) [5].

A recent Cochrane review [78] analyzed studies on the efficacy of pressure-relieving devices (mattresses and overlays) and came to the result that the use of pressure-relieving overlays during surgery reduced the postoperative incidence of pressure ulcers, although two studies reported adverse skin reactions after the use of foam mattresses or overlays.

A meta-analysis [79] confirmed the protective effect of pressure-relieving mattresses compared to standard mattresses, of high-specification foam mattresses compared to standard mattresses and of certain air-filled or foam overlays compared to standard mattresses with regard to the incidence of heel pressure ulcers. A detailed health technology assessment [80] and a careful systematic review [81] confirmed the efficacy of pressure-relieving overlays used on operating tables.

The superiority of these devices compared to standard care is evident. However, comparisons between different modes of action (devices) are lacking. Obese patients represent a separate patient cohort.

Positioning: The larger the area of the body being supported, the less pressure will be placed on tissue. In heel offloading, it is important to ensure that the heels are not elevated too high because this will increase pressure on the sacral area. The same applies if the head plate is too high. The operating table should be bent with the hip in a physiologically correct position. It is important to ensure the patient does not “slide down” (e.g. by placing a rolled towel under the buttocks) because sliding not only creates pressure but may also result in shearing.

Placing the heels on a soft underlay does not appear to be as effective as heel offloading, as a soft underlay will not necessarily prevent strong pressure being exerted on underlying deep tissue. Other positioning options include 30° oblique position, 135° angle sitting position, inclined position, etc. A systematic review showed that repositioning intervals could be reduced from every
2 hours to every 4 hours if a suitable pressure distribution device was used [82].

**Other measures:** According to a recent Cochrane review [83] the data from randomized controlled studies (RCTs) on the efficacy of prophylactic dressings (e.g. foam dressings or self-adhesive hydrocolloid dressings to relieve pressure) to reduce the incidence of pressure ulcers (e.g. on the heels) is still too limited to allow any recommendations to be made. The training of personnel as a supportive measure is often also studied and is recommended. Innovative techniques such as pressure measurement or the measurement of tissue perfusion at specific body sites (sacrum) are being evaluated in preclinical trials or in the form of additional monitoring which is compared to pressure-relieving devices. The efficacy (i.e., in reducing the incidence of pressure ulcers) of regular monitoring (with an alarm) in clinical practice has not yet been studied.

Recommendations for prevention (Fig. 8):
- Standardized risk assessments are not recommended.
- All patients should be treated as though they are at risk.
- The use of pressure-relieving overlays on operating tables is recommended.
- Ensure that the largest possible area of the patient is being supported.
- The prophylactic use of dressings (such as hydrocolloid dressing) to relieve pressure on healthy skin at specific sites at risk (over bony prominences) is not currently recommended.

### 6.5 Compartment syndrome

Acute compartment syndrome (CS) of the lower extremity is a particularly serious if rare form of positioning injury which has been almost exclusively described after lengthy operations with the patient in the lithotomy position [26,84–88]. CS refers to the resulting muscular and neuronal damage due to pathological increase in pressure within a confined inelastic space, in this case in one or more of the four muscular compartments of the lower leg [26,89]. If the diagnosis and start of treatment is delayed, consequences can be serious and may range from loss of function to amputation of the affected extremity and even the death of the patient from multi-organ failure. The reported incidence for gynecological procedures with the patient in the lithotomy position is reported to be between 0.028 and 0.28% depending on the patient cohort, although the estimated number due to diagnostic error and lack of attention appears to be higher [84,90,91]. This complication carries a high forensic relevance as treatment error is assumed to be causative in more than 50% of cases (Source: advisory body of the North Rhine Medical Association).

The precise etiology is unclear; however decreased perfusion pressure coupled with increased pressure on the tissue (caused by resting the calf on the support) appears to result in inadequate oxygen supply to tissues. This creates ischemia which in turn leads to a further increase in tissue pressure through the activation and release of mediators and toxic metabolic products and fluid leakage from vessels. If this vicious circle is not interrupted by fasciotomy to relieve the pressure, necrosis of affected structures can follow. Numerous experiments have shown that pressure drops significantly if the leg is positioned above the right atrium or the patient is placed in the Trendelenburg position [92]. An extreme lithotomy position reduces the median arterial pressure in the lower extremity to values which correspond to those measured for manifest CS, with this effect being reinforced in the Trendelenburg position [90,93,94]. These results suggest that minimizing the time during surgery in which the patient is placed either in a lithotomy or a Trendelenburg position is the best way of preventing CS. CS has been described after using a number of very different leg supports. Although the use of slings results in a lower application of pressure on the lower leg compared to other systems [92], routine use of slings is not recommended because of the associated higher rates of other neuropathies, particularly of peroneal neuropathies [95].

The diagnosis of CS is based on clinical examination. The first signs are usually pain at some distance from the surgical site (e.g. the leg) and diminished sensation and paresthesia in the affected area [96]. If an epidural catheter (EC) has been placed, the above-listed symptoms should not be exclusively ascribed to the EC; it is important to consider the possibility of incipient CS. One review found no delay in diagnosis despite postoperative analgesia if patients were adequately monitored [97].

To verify or exclude the possibility that sensory deficits and paresthesias are caused by analgesia administered in the vicinity of the spinal cord (e.g. EC), the anesthesiologist must be informed and consulted immediately. For the differential diagnosis it is useful to discontinue administration of the local analgesic through the catheter to check whether this is then followed by a drop in sensory and/or motor deficits.

Invasive measurement of intracompartmental pressure (ICP) provides additional information in cases where the constellation of symptoms is unclear, although the routine use of invasive measurement techniques is not indicated. The ICP threshold which indicates manifest CS is still controversially discussed in the literature [98]. The use of pulse oximeters to monitor pressure in the extremities offers no benefits as arterial perfusion and oxygen saturation of the extremities only drop at a very late stage [99].

To date, there are no evidence-based recommendations for prevention as none of the measures proposed in the literature have been validated in prospective studies. In a prospective observational study, a drop in the incidence of CS from 0.8 to 0 % was reported for a patient cohort at high risk of CS scheduled for extensive endometriosis surgery after implementing a combination of different measures [100]. These included minimizing the number of procedures performed with patients in the lithotomy position in favor of a modified supine position with the legs abducted, intermittent repositioning of the legs during surgery, and the use of vacuum mattresses to prevent the patient from slipping. The routine use of intermittent compression devices (mainly used in Anglo-American countries) has also been recommended; however other studies have reported an increased risk of complications associated with the use of these compression stockings.

Treatment of manifest CS consists of fasciotomy of all affected muscle compartments.

Recommendations for prevention:
- The time which the patient spends in the lithotomy position must be kept to a minimum (AST, AORN), particularly if access to the perineum or the vagina is not required. Use of other potential alternatives such as placing the legs flat at an abduction of ≤ 45° together with slight flexion of the hip should be considered.
- Where possible, the legs should be positioned at the level of or below the right atrium.
- The time spent by the patient in the Trendelenburg position should also be kept to a minimum; the patient should be moved from the Trendelenburg position and repositioned as soon as it is surgically possible.
Sliding of the patient in a cranial direction must be avoided using suitable positioning aids. None of the routinely used positioning devices (gel mat vs. vacuum mattress vs. foam underlay) have been shown to be superior to any of the alternatives.

The use of typical knee-and-lower-leg leg holders should be avoided. If such leg holders are used, additional padding with gel mats is necessary.

All medical professionals involved in the perioperative care and treatment of the patient must be aware of the possibility for CS and be familiar with the clinical signs of postoperative CS after lengthy operations performed with the patient in the lithotomy position.

Routine intraoperative measurement of the compartmental pressure for early diagnosis is not recommended because of the difficulty in defining the threshold values. Invasive pressure monitoring should be additionally performed for diagnosis if symptoms are unclear and there is a clinical suspicion.

Physiological rationale suggests that intermittent repositioning of the legs in operations of 3 hours can reduce intracompartmental pressure. There is currently no evidence that this will reduce compartmental pressure in the long term or prevent CS. Neither the repositioning intervals nor the appropriate duration of repositioning done intraoperatively have been validated in studies. Moreover, there is a potential risk that secondary positioning mistakes can occur due to unnoticed changes to the original (correct) position of the leg.

If the patient is additionally receiving analgesia administered in the vicinity of the spinal cord (e.g. epidural catheter), the anesthesiologist must be informed immediately. The anesthesiologist must make the differential diagnosis, identifying the causes of sensory and/or motor deficits and/or detecting a compartment syndrome where present.

7 Clinical Monitoring and Diagnosis of Positioning-related Injury

7.1 Intraoperative monitoring

Intraoperative monitoring is, by and large, unsuitable for detecting manifest positioning injury. In particular, neuropathies cannot be diagnosed in anesthetized patients. Instead, the main tasks of intraoperative monitoring are:

- controlling the patient’s position after planned repositioning; documenting the repositioning and its control; if leg holders and arm holder are used prior to repositioning the legs, ensuring that the patient’s fingers and hands do not get trapped or pinched;
- excluding unintentional critical changes to the (presumably) correct initial position of the patient.

Other than the recommendation that the patient’s position should be checked after planned repositioning, the start and the duration of the intervals for monitoring the patient’s position have not been defined.

The AORN guideline has suggested considering whether patients should be repositioned intraoperatively during lengthy surgical procedures (> 4 hours) to reduce the risk of pressure-related injury (pressure ulcers, neuropathies, CS) but does not make suggestions about the time and intervals for repositioning [2]. According to the most recent study reviews, there are no studies which show that active repositioning is effective for the prevention of positioning-related injury.

7.2 Postoperative monitoring

The likelihood of a positioning injury occurring can be minimized by carrying out various prophylactic measures described above; however, even if all possible care is taken, it is not possible to completely exclude positioning injury (s. also item 2.2) [15]. If an injury occurs, it is important to recognize this as early as possible, make the appropriate diagnosis and provide suitable care and treatment [17].

The goal of postoperative clinical monitoring must therefore be to detect positioning injuries early on, to provide early treatment where possible. Detailed early examination of the patient (for example EMG) may help to exclude the presence of preexisting neuropathies, which can be of great forensic importance.

At the first postoperative visit by the physician to the patient who is now conscious and responsive, it is important to ask specifically about the following symptoms or carry out a targeted examination, particularly after procedures performed in the lithotomy position which is associated with a higher risk of injury:

- reduced sensory perception and/or weakness in the extremities;
- diffuse pain with no immediate association to the direct surgical trauma;
- inspection of the skin on body areas at particular risk.

If there is a postoperative suspicion of neuropathy, it is important in addition to a potential positioning injury to consider other causes such as direct injury resulting from surgery; pressure injury resulting from hematoma or edema; and other previously unrecognized anatomical variants which can facilitate injury. A number of diseases such as diabetes mellitus, alcoholism, hypovitaminosis, uremia and malignant tumors are associated with a higher risk of neuropathies [20,101,102].

If specific symptoms are reported or findings of the detailed examination are suspicious, a neurological examination should be done promptly by another consultant for an objective assessment. In this context, physicians are referred to the S3-guideline on the care of peripheral nerve injuries: “Versorgung peripherer Nervenverletzungen” [32].

- Every patient who is operated on in the lithotomy position for > 3 hours should undergo a postoperative clinical examination (pain, motor function, sensory perception, measurement of extent and scope) and laboratory examination where necessary (determination of creatine kinase levels) for CS.

- Nursing staff must keep an eye on skin changes after surgical interventions, document any changes occurring and report them to the attending physician.

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