Acute compartment syndrome
Cause, diagnosis, and new viewpoint

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Abstract

Background: Acute compartment syndrome (ACS) is defined as a clinical entity originated from trauma or other conditions, and remains challenging to diagnose and treat effectively. The review was aim to present the controversy in diagnosing, treating ACS. It was found that there was no criterion about the ACS, and result unnecessary osteotomy. The presence of clinical assessment (5P) always means the necrosis of muscles and was the most serious or irreversible stage of ACS. Besides pressure methods, the threshold of pressure identifying ACS was also controversial.

Methods: Immediate surgical fasciotomy was important to prevent severe sequelae of the ACS. However, there was still controversy about the right time that fasciotomy should be done to avoid irreversible ischemic changes. The most important thing to treat ACS was comprehension to the true injury mechanism, but a systemic classification about traumatic mechanism in most literature was not clear.

Results: After observations to fracture patients with blister, we recommended that surgeons dealing with such emergencies should be vigilant, and the indication for fasciotomy should be strictly controlled following with injury mechanism especially for patients without severe soft tissue injury.

Conclusion: For those crushing and soft tissue injuries, the current evidence based strategies for managing patients was useful, but for those fracture related injury, more examination was necessary to avoid overtreatment especially for those patients with blister observed. In facing patients, medical history, injured mechanism should be paid special attention, and rigorous classification about traumatic etiology was the key for the treatment of these patients.

Abbreviation: ACS = acute compartment syndrome.

Keywords: acute compartment syndrome, diagnosis, fasciotomy, mechanism

1. Introduction

Acute compartment syndrome (ACS) is defined as a clinical entity originated from trauma or other conditions that cause bleeding, edema, or that compromises perfusion in limbs. It was described around 130 years ago, and remains challenging to diagnose and treat effectively.\textsuperscript{1} After a decrease in a compartment volume or and an increase in the contents of a compartment, ACS occurs when pressure increases within a confined closed fascial space causing subsequently reduced blood flow and tissue perfusion that may lead to ischemic pain, possible soft-tissue damage, and limb-threatening. With a character of increased intracompartmental pressure, it is a surgical emergency and commonly occurs in the lower leg, forearm, thigh, foot, gluteal region, hand, and abdomen. The incidence rate of ACS was reported 30.4% especially in shaft and proximal regions of tibia. The higher incidence of ACS in proximal tibia is directly related to high energy trauma causing comminuted fractures, especially those affecting the entity of medial tibial plateau and increased with the fracture line extended laterally.\textsuperscript{2–5}

To prevent serious complications induced by ACS, fasciotomy should be done before irreversible tissue necrosis occurs, thus there is a strong clinic bias toward doing fasciotomy empirically or prophylactically in patients who are considered to be at high risk and/or who have concerning clinical findings. It is generally recommended that conducting a prophylactic fasciotomy, which may be unnecessary, is better than do it too late, or ignoring a true ACS, given the potential systemic risks (rhabdomyolysis and myonecrosis) and functional loss associated with untreated ACS (Supplemental Figure 1, http://links.lww.com/MD/D60). However, there were controversies regarding the diagnosis (method and pressure threshold), treatment (the time of fasciotomy), and injury mechanism of ACS. The aim of this review was to present the uncertainties in treating ACS and propose new solutions.
2. Diagnosis varieties

The patients’ photographs were collected from our institutional database, retrospectively, and all the methods were performed in accordance with the relevant guidelines and regulations of the Third Hospital of Hebei Medical University. It was approved by the committee of the Third Hospital of Hebei Medical University.

The diagnosis of compartment syndrome is always controversial and is based on clinical assessment and pressure measurement in compartment. Compartment syndrome clinical hallmarks have been defined as the 5Ps: pain out of proportion, pallor, paresthesia, paralysis, and pulselessness. Someone considered 5Ps as the established diagnostic procedures monitoring ACS, but it is an unreliable determinant of the presence of the syndrome, and many of clinical symptoms and signs also occur in patients without ACS (reamed nailing to tibia fractures with a sign of common or deep peroneal nerve), perhaps due to direct tissue injury 10 to 18. It is also not useful in patients with a decreased level of consciousness, unconscious, or insensate who are unable to provide feedback. In fact, the presence of these signs always means the necrosis of muscles and was the most serious or irreversible stage of ACS. Furthermore, these signs are more often signs of arterial ischemia than ACS, and be elicited only in the fully alert patient.[6–8]

The surgeons concern mainly about clinical signs of ACS such as worsening pain that is out of proportion and increasing analgesic requirements, or anxiety, agitation. However, the subjective symptoms rely heavily on clinical expertise, are impossible to standardize, and meanwhile the early changes of ischemia in compartment was the increased pressure.[9] Ulmer found that clinical examination had poor sensitivity and a high negative predictive value, which means that it is better in excluding than confirming ACS.[10] Therefore, subjective clinical assessments of compartments are unreliable even judged by experienced clinicians, the results was that unnecessary liberal fasciotomy which expose patients to an increased risk of complications from wound infection was more than we can image. In the case that the clinical diagnosis is equivocal, measurement of intracompartmental tissue pressure might be helpful because of the fact that the pressure changes pull ahead the clinical symptoms and signs.[10] Through pressure monitoring, it reduces both the delay to fasciotomy and the development of sequelae, but the researches that identified the pressure as the measuring criteria was only 11.7%.[11] An investigation reveals that only about half of the hospitals in Germany have the technical equipment to do this despite most surgeons’ agreeing that pressure measurement is the standard method.

There are many methods of monitoring pressure, and the equipment was different from each other. Invasive choices of monitoring pressure include the Whitesides needle manometer, a slit or wick catheter. The deficiency of these devices was that it was easily to be blocked by luer muscle and blood clots. The STIC Monitor (Stryker Orthopaedics, Mahwah, NJ) is a portable monitor that uses a side port needle, a disposable syringe of saline flush, and a digital read out manometer to allow for simple measurement of compartment pressure. The measurement will be effected by position, measuring location and tissue in tip of needle. Through knee cadaveric specimens, large calculated interobserver technical variations, and errors in the measurement of compartment pressures, he found that even with proper technique, 40% of the measurements were > 5 mm Hg from the actual pressure. Therefore, it was misleading to make the choice depending on pressure monitor; many patients may be wrongly diagnosed as ACS. Some researches considered that the pressure monitor by needle was technically wrong in clinic. Nudel et al found that ACS is not uniform about pressure distribution in the compartment.[6] The pressure adjacent to the damaged artery is substantially higher than a critical value, indicating the need for fasciotomy. However, at the same time, the pressure in regions far from the bleeding artery is substantially lower than the threshold value even after 2 hours from injury. Intracompartmental pressure was highest and should be measured within 5 cm of the site of fracture.[12,13] However, the standard whether one should obtain pressures near the fracture, or measure further away (outside the zone of injury) to obtain a more representative pressure to the majority of the compartment was not established.

Besides pressure methods, the threshold of pressure identifying ACS was also controversial. It was identified a compartment pressure of 30 mm Hg was resistant to infusion of fluid and recommended as indication of surgical fasciotomy.[14] Levels ranging from 30 to 50 mm Hg are also proposed as critical level to diagnose ACS.[14–16] However, the value was proved to be inaccurate, and no correlation was found between high compartment pressure and clinical outcomes in patients treated primarily in plaster cast.[17] More and more researches demonstrated that pressures more than 30 mm Hg can also be tolerated without sequel, and proposed that instead of an absolute threshold, the difference between diastolic pressure and compartment pressure was accepted as indication for fasciotomy (diastolic pressure minus compartment pressure <30 mm Hg).[18–21] However, along with pressure measurement, it is important that the diagnosis of ACS should take into account time factors, and single pressure measurements alone reflect instantaneous blood perfusion. The tissue and blood pressure was dynamic equilibrium, and pressure in different compartment was various from each other. Bussell et al reported that pressure in the anterior compartment was higher compared to all the other compartments within the healthy and fractured leg in children.[22] McQueen et al monitored pressure for 2 hours and demonstrated excellent sensitivity (94%) and specificity for ACS after tibia fractures.[23] On the contrary, there was also researches reported that continuous compartment pressure monitoring was not advocated in alert patients.[24] Diagnosis based on pressure measurements (diastolic pressure minus compartment pressure <30 mm Hg) alone is also reported unreliable. Ho found high compartment pressures are frequently seen in patients with tibial shaft fractures, but in most cases, it does not equate to the presence of compartment syndrome, and unnecessary fasciotomy should be avoided.[25] Pryason et al followed blood pressure and compartment pressure in 19 patients with lower extremity fractures, and also did not found compartment syndrome with the perfusion pressure threshold (<30 mm Hg).[26] Janzing and Broos reported that 45.4% of patients would have undergone fasciotomies using ΔP under 30 mm Hg, leading to a number of unnecessary operations and complications.[27]
The incidence of complications associated with fasciotomy has been reported to be as high as 87% following battlefield trauma.\cite{28} In earthquake, fasciotomy was always carried out without pressure measurement.\cite{29}

Above all, when the clinical observations are inconclusive, pressure measurement can be helpful to confirm or exclude the diagnosis, not as a screening tool for those with an increased risk of developing compartment syndrome. However, there was no standard criterion even when clinical indication and pressure threshold combined that enable a successful prediction of the need for surgery treatment. And after considering the confusion about ACS, we begin to doubt the accuracy of ACS, and the existence of ACS.

### 3. The time of fasciotomy and results

As we mentioned earlier, in traditional view, immediate surgical fasciotomy was important to prevent severe suquelea of the ACS. However, there was still controversy about the right time that fasciotomy should be done to avoid irreversible ischemic changes.

The ischemic necrosis of muscle can be observed as early as 3 hours, 5% may be injured after 4 hours, and become permanent in 8 hours.\cite{30,31} Labbe et al reported that when the ischemia was 3, 4, and 3 hours, the necrosis of leg muscles was up to 20%, 30%, and 90%, respectively.\cite{32} On the contrary, Sheridan and Matsen found that 68% of 22 patients treated within 12 hours recovered normal lower extremity function compared to only 8% treated after 12 hours.\cite{33} There was not only damage to muscles but also to the associated nerves traversing the compartment if compartment syndrome is not diagnosed and treated early. It was reported that if compartmental release was performed within 4 hours, the nerve conduction velocity returned to normal regardless of the amount of pressure or time of the pressure applied. However, nerve conduction velocity will not return to normal if the release was performed after 12 hours. The late diagnosis may result in the possibility of irreversible nerve, muscle damage, amputation, and even death.

Despite there is obvious evidence that delay in treatment leads to poorer outcomes, it is difficult to determine the exact time of performance for fasciotomy. For delayed compartment syndrome in adults, someone propose that routine fasciotomy should be conducted to prevent more morbidity and complication.\cite{34} In contrast, others suggested that 8 hours of ischemia can result in permanent myonecrosis, so traditional fasciotomies should be restricted as fasciotomies can no longer reverse the nerve damage and muscle necrosis, and the procedure may potentially lead to wound problems and infection. Reis and Michaelson suggested that the skin serves an important role as a barrier to infection, and wound problems and infection. Reis and Michaelson suggested that the skin serves an important role as a barrier to infection, and wound problems and infection.

Finkbeiner et al also reported that after delayed fasciotomy has greater morbidity than the muscle contractures occurred from myonecrosis.\cite{36} Ritenour et al reported that wounded soldiers that had delayed fasciotomies, had higher rates of muscle excision, amputation, and mortality.\cite{37} Therefore, prophylactic fasciotomy can also induce major complications, and the risk–benefit ratio should be weighed heavily.

However, functional outcomes were different and controversial in different aged patients. Livingston et al reported the time from symptom onset or initial injury to ACS diagnosis in nonfracture pediatric patients to be over 48 hours.\cite{38} After a system review, Lin and Balch Samora reported that pediatric patients could still achieve functional recovery in 24 hours.\cite{14} Kanj et al found that although ACS of the upper extremity in children is often associated with a long delay between injury and fasciotomy, most children still achieve excellent outcomes.\cite{19} Above all, increased intracompartamental pressure can be tolerated for longer time compared with adults before tissue necrosis becomes irreversible. There was more time allowed to diagnose pediatric ACS after identifying the exact mechanism of injury.

### 4. Injury mechanism and fasciotomy results

The consequences of missed diagnosis are severe for both patients and surgeons. It is important to be cognizant that there is no universal pressure measurement that serves as the threshold for fasciotomy. The main etiologies of ACS was traumatic injuries such as fracture and crush-type injury, while other injuries such as limb ischemia (ischemia-reperfusion injury after revascularization), tourniquet, tight splint, shock trousers, drug injection, or snake bites could also induce ACS.\cite{30} Furthermore, the most important thing to treat ACS was comprehension to the true injury mechanism, but a systemic classification about traumatic mechanism in most literature was not clear. Patients in most articles can be classified into soft-tissue injury related, vascular injury related, fracture related according to the injured anatomical structure.\cite{41}

Soft-tissue injury can be considered as crush syndrome. Patients with crush syndrome are induced by continuous prolonged pressure on muscle tissues, and character with massively swollen limbs, shock, myoglobinuria, and renal failure. Although fasciotomy is considered as the gold standard for ACS, but the role of fasciotomy in the treatment of crush syndrome is still controversial. The most debate was that in one hand, early fasciotomy prevents further muscle damage, and there were researches recommend fasciotomy for patients with crush syndrome.\cite{29,42,43} On the other hand, the fasciotomy convert a closed wound into an open one which increases the rate of infection. Despite high or increasing intracompartamental pressure, conservative treatment was advocated unless open wound existed or the limb circulation is decreased.\cite{33,44-46} For those patients with the necrotic muscle, Huang et al suggested that fasciotomy offers no benefit but increases the rate of infection and amputation.\cite{45}

The ACS was always found to be mixed with the concept of crushing syndrome and Volkmann contracture. Crush syndrome is a medical condition that can be caused by a “crush injury,” and skeletal muscle becomes damaged under the heavy weight. The injury mechanism includes earthquakes, traffic accident, and war conflict and so on. The saved entrapped victims without immediate medical treatment caused the damage to the structure of compartment and lead muscle necrotic or damaged, and induce acute kidney injury that requires rapid and special treatment. The early debridement (not fasciotomy) should only be considered in patients with crushing injuries. Vascular injury associated with orthopedic trauma is also a potentially limb- and life-threatening, if the diagnosis of an associated vascular injury is missed or delayed, Volkmann contracture presented in late status (Table 1).

Reperfusion was one reason reported for the development of ACS, and different from complete ischemia, ACS causes myonecrosis in the face of patent vessels 68. Furthermore, there
Fracture blisters were reported as a relatively uncommon complication of high-energy fractures (2.9%), but most tibial fracture can be observed with clear fluid or blood filled blister as early as 6 hours postfracture. It was considered that the factor in development of blister is injury to the dermal-epidermal junction resulting from high shear in the skin. The retention of some degree of epidermal cells in the clear-filled blisters contributed to a faster reepithelialization compared with blood-filled blister. However, the theory cannot explain the blister remote from the fracture deformity. Halawi also found that the blister which may induce infection can be observed after primary total knee arthroplasty, and the etiology of blister was multifactorial. The structure of skin was actually intact and without dermal disturbance, debridement of all devitalized tissue should be performed. Following experiment will be conducted soon to verify the hypothesis.

5. Conclusion

The ACS is considered as an orthopedic emergency which can lead to limb and life-threatening outcome if there is delay in diagnosis and treatment. Surgeons that involved in dealing with such emergencies should be vigilant, and the indication for fasciotomy should be strictly controlled following with injury mechanism especially for patients without severe soft-tissue injury. For those crushing and soft-tissue injuries, the current evidence-based strategies for managing patients were useful, but for those fracture-related injury, more examination was necessary to avoid overtreatment especially for those patients with blister observed. In facing patients' medical history, injured mechanism should be paid special attention, and rigorous classification about traumatic etiology was the key for the treatment of these patients.

Author contributions

Zhiyong Hou designed the study. Jialiang and Yingchao Yin conducted the procedure and collected the data of the study.
Jiali Guo, Lin Jin and Ruipeng Zhang drafted the manuscript. Zhiyong Hou and Yingze Zhang revised and approved the final manuscript.

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References

[1] von Volkmann R. Ischaemic muscle paralyses and contractures. 1881. Clin Orthop Relat Res 2007;456:20–1.
[2] Acklin YP, Potocnik P, Sommer C. Compartment syndrome in dislocation and non-dislocation type proximal tibia fractures: analysis of 356 consecutive cases. Arch Orthop Trauma Surg 2012;132:227–31.
[3] Ziran BH, Becher SJ. Radiographic predictors of compartment syndrome in tibial plateau fractures. J Orthop Trauma 2013;27:612–5.
[4] Chang YH, Yu YK, Yeh WL, et al. Tibial plateau fracture with compartment syndrome: a complication of higher incidence in Taiwan. Chang Gung Med J 2002;23:149–53.
[5] Stark E, Stucken C, Trainer G, et al. Compartment syndrome in Schatzker type VI plateau fractures and medial condylar fracture-dislocations treated with temporary external fixation. J Orthop Trauma 2009;23:502–6.
[6] Nudel I, Dorfmann L, Debotton G. The compartment syndrome: is the intra-compartment pressure a reliable indicator for early diagnosis? Math Med Biol 2017;34:547–58.
[7] Ulmer T. The clinical diagnosis of compartment syndrome of the lower leg: are clinical findings predictive of the disorder? J Orthop Trauma 2002;16:572–7.
[8] Shuler FD, Dietz MJ. Physicians’ ability to manually detect isolated elevations in leg intracompartmental pressure. J Bone Joint Surg Am 2010;92:361–7.
[9] Bae DS, Kadiyala RK, Waters PM. Acute compartment syndrome in children: contemporary diagnosis, treatment, and outcome. J Pediatr Orthop 2001;21:680–8.
[10] McQueen MM, Christie J, Court-Brown CM. Compartment pressures after intramedullary nailing of the tibia. J Bone Joint Surg Br 1990;72:395–7.
[11] O’Toole RV, Whitney A, Merchant N, et al. Variation in diagnosis of compartment syndrome by surgeons treating tibial shaft fractures. J Trauma 2009;67:735–41.
[12] Beckman MM, Whitesides TE Jr, Grewe SR, et al. Compartment pressure in association with closed tibial fractures. The relationship between tissue pressure, compartment, and the distance from the site of the fracture. J Bone Joint Surg Am 1994;76:1285–92.
[13] Matava MJ, Whitesides TE Jr, Seiler JG3rd, et al. Determination of the compartment pressure threshold of muscle ischemia in a canine model. J Trauma 1994;37:30–8.
[14] Mubarak SJ, Owen CA, Hazengs AR, et al. Acute compartment syndromes: diagnosis and treatment with the aid of the wick catheter. J Bone Joint Surg Am 1978;60:1091–5.
[15] Allen MJ, Stirling AJ, Crawshaw CV, et al. Intracompartmental pressure monitoring of leg injuries. An aid to management. J Bone Joint Surg Br 1985;67:653–7.
[16] Mubarak SJ, Hazengs AR. Acute compartment syndromes. Surg Clin N Am 1983;6:539–65.
[17] Triffitt PD, Konig D, Harper WM, et al. Compartment pressures after closed tibial shaft fractures. Their relation to functional outcome. J Bone Joint Surg Br 1992;74:195–8.
[18] Shadgan B, Menon M, O’Brien PJ, et al. Diagnostic techniques in acute compartment syndrome of the leg. J Orthop Trauma 2008;22:581–7.
[19] Elliott KG, Johnstone AJ. Diagnosing acute compartment syndrome. J Bone Joint Surg Br 2003;85:625–32.
[20] White TO, Howell GE, Will EM, et al. Elevated intramuscular compartment pressures do not influence outcome after tibial fracture. J Trauma 2003;55:1133–8.
[21] McQueen MM, Court-Brown CM. Compartment monitoring in tibial fractures. The pressure threshold for decompression. J Bone Joint Surg Br 1996;78:99–104.
[22] Russell HR, Aufdenblatten CA, Subotic U, et al. Compartment pressures in children with normal and fractured lower extremities. Eur J Trauma Emerg Surg 2019;4: doi: 10.1007/s10199-019-01082-9.
[23] McQueen MM, Christie J, Court-Brown CM. Acute compartment syndrome in tibial diaphyseal fractures. J Bone Joint Surg Br 1996;78:95–8.
[24] Harris IA, Kadir A, Donald G. Continuous compartment pressure monitoring for tibia fractures: does it influence outcome? J Trauma 2006;60:1330–5.
[25] Lor KKH, Yeoh NCS, Wong KP, et al. Raised compartment pressure frequencies are frequently observed with tibial shaft fractures despite the absence of compartment syndrome: a prospective cohort study. J Orthop Surg (Hong Kong) 2017;25:2309499017717362 doi: 10.1177/2309499017717362.
[26] Prayson MJ, Chen JL, Hampers D, et al. Baseline compartment pressure measurements in isolated lower extremity fractures without clinical compartment syndrome. J Trauma 2006;60:1347–54.
[27] Kanj WW, Gunderson MA, Carrigan RB, et al. Acute compartment syndrome: a review of 39 cases. J Pediatr Orthop 2008;64:S153–61.
[28] Mubarak SJ, Owen CA. Compartment syndrome and sequential complications. BMC Musculoskelet Disord 2019;20:98.
[29] Brien PJ, et al. Diagnostic techniques in acute compartment syndrome of the leg: are clinical findings predictive of the disorder? J Orthop Trauma 2002;16:572–7.
[30] Medscape Reference. Fasciitis caused by the Hanshin-Awaji earthquake. J Trauma 1997;47:2470–5.
[31] Whitesides TE, Beckman MM. Acute compartment syndrome: update on diagnosis and treatment. J Am Acad Orthop Surg 1996;4:209–18.
[32] Beckman MM, Whitesides TE Jr, Grewe SR, et al. Histologic determination of the ischemic threshold of muscle in the canine compartment syndrome model. J Orthop Trauma 1993;7:199–210.
[33] Kurihara H, Tajiri H, Takamatsu Y, et al. Incidence of decompression before the 1995 Hanshin-Awaji earthquake. J Trauma 1997;43:516–21.
[34] Toole RV, Whitney A, Merchant N, et al. Variation in diagnosis of compartment syndrome by surgeons treating tibial shaft fractures. J Trauma 2009;67:735–41.
[35] Heiner R, Dau N, Winkler M, et al. Compartment syndrome: a complication of higher incidence in Taiwan. Chang Gung Med J 2002;23:149–53.
[36] Stark E, Stucken C, Trainer G, et al. Compartment syndrome in Schatzker type VI plateau fractures and medial condylar fracture-dislocations treated with temporary external fixation. J Orthop Trauma 2009;23:502–6.
[37] Nudel I, Dorfmann L, Debotton G. The compartment syndrome: is the intra-compartment pressure a reliable indicator for early diagnosis? Math Med Biol 2017;34:547–58.
[38] Ulmer T. The clinical diagnosis of compartment syndrome of the lower leg: are clinical findings predictive of the disorder? J Orthop Trauma 2002;16:572–7.
[39] Shuler FD, Dietz MJ. Physicians’ ability to manually detect isolated elevations in leg intracompartmental pressure. J Bone Joint Surg Am 2010;92:361–7.
[40] Bae DS, Kadiyala RK, Waters PM. Acute compartment syndrome in children: contemporary diagnosis, treatment, and outcome. J Pediatr Orthop 2001;21:680–8.
[41] McQueen MM, Christie J, Court-Brown CM. Compartment pressures after intramedullary nailing of the tibia. J Bone Joint Surg Br 1990;72:395–7.
[42] O’Toole RV, Whitney A, Merchant N, et al. Variation in diagnosis of compartment syndrome by surgeons treating tibial shaft fractures. J Trauma 2009;67:735–41.
[43] Beckman MM, Whitesides TE Jr, Grewe SR, et al. Compartment pressure in association with closed tibial fractures. The relationship between tissue pressure, compartment, and the distance from the site of the fracture. J Bone Joint Surg Am 1994;76:1285–92.
[44] Matava MJ, Whitesides TE Jr, Seiler JG3rd, et al. Determination of the compartment pressure threshold of muscle ischemia in a canine model. J Trauma 1994;37:30–8.
[45] Mubarak SJ, Owen CA, Hazengs AR, et al. Acute compartment syndromes: diagnosis and treatment with the aid of the wick catheter. J Bone Joint Surg Am 1978;60:1091–5.
[46] Allen MJ, Stirling AJ, Crawshaw CV, et al. Intracompartmental pressure monitoring of leg injuries. An aid to management. J Bone Joint Surg Br 1985;67:653–7.
[47] Mubarak SJ, Hazengs AR. Acute compartment syndromes. Surg Clin N Am 1983;6:539–65.
[48] Triffitt PD, Konig D, Harper WM, et al. Compartment pressures after closed tibial shaft fracture. Their relation to functional outcome. J Bone Joint Surg Br 1992;74:195–8.
[49] Shadgan B, Menon M, O’Brien PJ, et al. Diagnostic techniques in acute compartment syndrome of the leg. J Orthop Trauma 2008;22:581–7.
[50] Elliott KG, Johnstone AJ. Diagnosing acute compartment syndrome. J Bone Joint Surg Br 2003;85:625–32.
[51] White TO, Howell GE, Will EM, et al. Elevated intramuscular compartment pressures do not influence outcome after tibial fracture. J Trauma 2003;55:1133–8.
[47] Huang KC, Lee TS, Lin YM, et al. Clinical features and outcome of crush syndrome caused by the Chi-Chi earthquake. J Formos Med Assoc 2002;101:249–56.

[48] Yeon J, Jung YW, Yang SS, et al. Lower limb compartment syndrome by reperfusion injury after treatment of arterial thrombosis post-laparoscopic radical hysterectomy and pelvic lymph node dissection for cervical cancer. Obstet Gynecol Sci 2017;60:223–6.

[49] Feliciano DV, Cruse PA, Spjut-Patrinely V, et al. Fasciotomy after trauma to the extremities. Am J Surg 1988;156:533–6.

[50] Lim LT, Michuda MS, Flanigan DP, et al. Popliteal artery trauma. 31 consecutive cases without amputation. Arch Surg 1980;115:1307–13.

[51] Rolls DL, Bernhard VM, Towne JK. Fasciotomy: an appraisal of controversial issues. Arch Surg 1981;116:1474–81.

[52] Abouezzi Z, Nassoura Z, Ivatury RR, et al. A critical reappraisal of indications for fasciotomy after extremity vascular trauma. Arch Surg 1998;133:547–51.

[53] Bermudez K, Knudson MM, Morabito D, et al. Fasciotomy, chronic venous insufficiency, and the calf muscle pump. Arch Surg 1998;133:1356–61.

[54] Olson SA, Glasgow RR. Acute compartment syndrome in lower extremity musculoskeletal trauma. J Am Acad Orthop Surg 2005;13:436–44.

[55] McQueen MM, Gaston P, Court-Brown CM. Acute compartment syndrome. Who is at risk? J Bone Joint Surg Br 2000;82:200–3.

[56] Patel RV, Haddad FS. Compartment syndromes. Br J Hosp Med 2005;66:583–6.

[57] Giordano CP, Scott D, Koval KJ, et al. Fracture blister formation: a laboratory study. J Trauma 1995;38:907–9.

[58] Halawi MJ. Fracture blisters after primary total knee arthroplasty. Am J Orthop 2015;44:E291–3.

[59] Varela CD, Vaughan TK, Carr JR, et al. Fracture blisters: clinical and pathological aspects. J Orthop Trauma 1993;7:417–27.