

KNEE-ANKLE-FOOT ORTHOSES FOR TREATING POSTERIOR KNEE PAIN RESULTING FROM GENU RECURVATUM: EFFICIENCY, PATIENTS' TOLERANCE AND SATISFACTION

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Objective: To assess the efficiency of knee-ankle-foot orthoses for treating painful genu recurvatum, and to determine users' tolerance and satisfaction.

Patients: Patients included in the study had a genu recurvatum during the stance phase, confirmed by a medical doctor on physical examination. A total of 27 patients with 31 knee-ankle-foot orthoses were included.

Methods: The main outcome was scored on a verbal numerical rating scale (VNRS) before and at least 3 months after a knee-ankle-foot orthosis was fitted, and scored on a verbal numerical pain rating scale (VRS). Secondary outcomes were rated with the Quebec User Evaluation of Satisfaction with assistive Technology (QUEST).

Results: After fitting the knee-ankle-foot orthosis, the median VNRS pain score decreased from 85/100 to 25/100 ($p \leq 0.001$) and the description of pain on the VRS decreased from "extreme" to "mild" ($p \leq 0.001$). The QUEST total score was 4.0.

Conclusion: Treating a painful genu recurvatum with a knee-ankle-foot orthosis reduced the pain efficiently whatever the patients' diagnosis, and high scores were obtained for patients' satisfaction.

Key words: knee-ankle-foot orthosis; KAFO; genu recurvatum; painful recurvatum.

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Genu recurvatum (GR) is hyperextension of the knee of more than 5°, in which the ground reaction force (GRF) line is anterior to the knee axis (1). GR can occur in various neuromuscular and musculoskeletal diseases, such as myopathy, upper motor neurone diseases, such as stroke, cerebral palsy, multiple sclerosis (2–5), and lower motor neurone diseases, such as poliomyelitis. GR is relatively common in stroke patients, where its rate of occurrence ranges from 40% to 68% (6).

The main physiopathological mechanisms of GR are weakness of the knee extensor, knee extensor spasticity, especially of the quadriceps, weakness of the buttock muscles leading to forward pelvic tilt, lumbar hyper-

lordosis and excessive hip flexion compensated by knee hyperextension, weakness of the knee flexors, limited dorsiflexion of the ankle with knee hyperextension due to the patients' inability to move the tibia forward during the stance phase, avoidance of painful pressure on the ball of the foot or proprioceptive disorders (5, 7).

Depending on the aetiology of GR, treatments include physical therapy, functional methods of electrical stimulation or electrogoniometric feedback (8, 9), botulinum toxin A injections in cases in which GR is due to spasticity (10, 11), orthoses, such as ankle-foot orthoses (AFOs) (12, 13), knee-ankle-foot orthoses (KAFOs) (1), knee orthoses (Swedish knee cages) (14) or hinged soft knee orthoses (15), orthopaedic surgery (16), or selective tibial neurectomy (17). Making clinical decisions in GR is difficult because of the wide range of aetiologies. Step-by-step analysis can be useful to define the main cause of GR and choose the best treatment (18). Studies on orthotic devices for GR have been conducted on AFO and KAFO users (1), the use of knee orthoses (Swedish knee cages) has been assessed (19), and the effectiveness of AFOs vs KAFOs has been compared (20, 21). To assess the effectiveness of KAFO treatment, some authors have analysed the improvement in gait parameters (20–24), while others have studied energy savings (13, 25–28).

In our clinical practice, posterior knee pain resulting from a GR, which is a frequent reason for consultation, consists of pain in the posterior structures of the knee associated with abnormal stretching of the posterior joint capsule and posterior knee ligaments. When repetitive loading occurs, posterior joint laxity can develop as a result of the tensile forces exerted on the posterior capsule, generating posterior articular, muscular and tendon pain (5, 7, 29–31), which can lead to a KAFO being fitted in order to restrict the GR and reduce the pain. Depending on the aetiology and cause of GR, a KAFO with a free knee joint and a posterior offset can be prescribed to reduce the pain. Several studies have described the effects of orthoses on gait parameters (14, 21, 22, 32), but no studies have focused on their effects in terms of pain relief.

The aim of this study was to assess the efficiency of KAFOs for treating painful GR and to determine users' tolerance and satisfaction.

MATERIALS AND METHODS

Experimental methods

This retrospective study was conducted on patients who had been fitted with KAFOs in order to alleviate posterior knee pain resulting from GR at our outpatient department of Physical and Rehabilitation Medicine. Data used were collected by an investigator who was not involved in the subjects' treatment.

Participants provided informed verbal consent after being given information about the study.

Population

Inclusion criteria. Patients with painful GR were included in the study. GR was defined as abnormal hyperextension of more than 5° (measured with a goniometer at physical examination) and by a ground reaction force line anterior to the knee axis during the stance phase (with the hip in the anatomical position). The indication for a KAFO was pain and not instability. Patients included in this study had to be at least 18 years of age and able to understand and answer the questionnaire. Patients with cognitive disorders, particularly memory disorders, were excluded. Additional inclusion criteria were a gait perimeter of more than 10 m without the KAFO, and having worn the orthosis for more than 3 months since the last correction.

Primary and secondary outcomes

Assessment was performed after the KAFO was fitted, and had been worn for at least 3 months, because of the retrospective nature of the study. Data were collected through patient questionnaires.

Primary outcomes. *Knee pain, before and after having the KAFO fitted, was measured with:*

- a verbal numerical pain rating scale (VNRS), giving a score out of 100 (where 0 means "no pain at all" and 100 means "worst imaginable pain").
- a verbal rating scale (VRS), where each response option consisted of descriptions of pain. Scores were attributed to the various responses corresponding to different levels of pain intensity: "no pain" [0], "mild pain" [1], "moderate pain" [2], "severe pain" [3], and "extreme pain" [4] (33).

The patient may also have had other pain localizations, but we only consider pain related to wearing a KAFO. Because of the retrospective nature of the study knee pain without the KAFO was evaluated by patients when they walked without the KAFO, even if they had already been fitted with a KAFO.

Secondary outcomes

The Quebec User Evaluation of Satisfaction with assistive Technology (QUEST) was used to measure patient satisfaction with the orthosis. This questionnaire comprises 12 satisfaction items (8 questions about the orthotic device, its dimensions, its weight, whether it was easy to adjust, its safety and security, its durability, whether it was easy to use, comfort, and effectiveness, and 4 questions about the services involved: its delivery, repairs and maintenance, and the professional services and follow up services), where all the responses to these items give a quantitative score (19, 34), as follows: 1: "not satisfied at all"; 2: "not very satisfied"; 3: "more or less satisfied"; 4: "quite satisfied"; and 5: "very satisfied".

The following 5 subjective gait parameters were rated without and with the KAFO: "much improved" [+2], "improved" [+1], "no difference" [0], "worse" [-1], "much worse" [-2] (35, 36) and scored only by patients:

- walking distance
- walking speed
- a sensation of recurvatum when walking
- self-confidence when walking
- feeling safe when walking.

Tolerance of the KAFO was assessed as follows:

- length of time spent wearing the orthosis per day.
- which activities of daily living could be carried out while wearing the orthosis, such as self-care (washing oneself, caring for body parts, toileting, dressing, looking after one's health) mobility (walking, lifting and carrying objects, using transportation, driving) and domestic life (acquisition of goods and services, preparation of meals, doing housework).

We asked patients about falls before and after delivery of the orthosis: ("falls: yes/no").

Statistical analysis

Paired Wilcoxon tests were used to analyse differences between the pain assessments (in terms of the VNRS and VRS scores) before and after a KAFO was fitted. McNemar's test was used to compare the proportion of fallers before and after the KAFOs were fitted. To calculate the results of the QUEST, the original methods described in the study by Demers et al. were used (34, 37) with means and standard deviation (SD).

To analyse the quantitative scores, such as the numerical verbal satisfaction scores ranging from 1 to 100, the median, quartiles and range were calculated. In the case of discrete characteristics with available meanings, such as the subjective assessment of walking without and with the KAFO (ranging from +2, "much improved", to -2, "much worse"), median values, quartiles and range were calculated.

An alpha £0.05 was taken to be statistically significant. All statistical analyses were performed using IBM SPSS Statistics 20.0 software.

RESULTS

Study population

Based on patients' files recorded between 2007 and 17 October 2014, 216 had a KAFO and 104 had a KAFO to treat GR. In this group of 104 patients, 33 had a KAFO because of a painful GR. Two patients declined to participate in the study, and 4 gave up wearing the KAFO before the end of the first month and declined to participate in the study (Fig. 1).

Patients' characteristics. The patients' characteristics and the types of KAFO are shown in Table I.

Diagnoses and clinical examination data. The patients' diagnoses, range of motion, and muscular strength are shown in Tables II and III.

KAFO characteristics. Twenty-five KAFOs were prescribed with a free offset knee joint, but a posterior

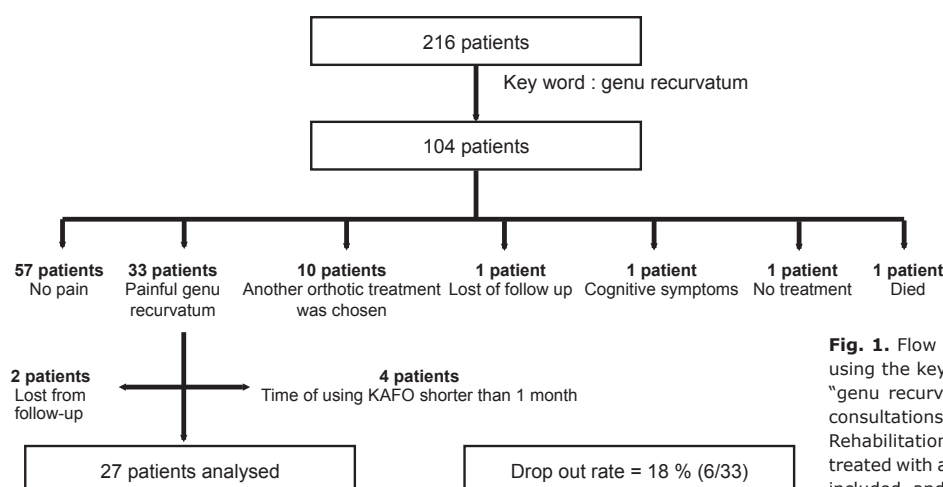


Fig. 1. Flow chart of the study population selected using the key words “knee-ankle-foot orthosis” and “genu recurvatum” from all the patients attending consultations in our Department of Physical and Rehabilitation Medicine. Only patients who were treated with a knee-ankle-foot orthosis for pain were included, and instability was not taken into account.

Table I. Characteristics of the patients’ in the study population ($n = 27$)

	Mean	Range
Age, years	56	(30–81)
Sex (F/M), n (%)	20 (74)/7 (26)	
Weight, kg	63	(43–84)
Height, m	1.64	(1.50–1.82)
Body mass index, kg/m^2	23.5	(16.8–31.6)
Mean GR measured on a table ($^\circ$)	18	(5–35)
Number of KAFOs	31:	
	11 on left side	
	12 on right side	
	4 on both sides	
Type of knee joint	25 with a free knee (80%) with posterior offset	
	6 with a locked knee (20%)	
Age of the current KAFOs	3 years	(4 months to 7 years)
How long patients had been wearing KAFOs	For 9 years	(4 months to 39 years)

F: female; M: male; KAFO: knee-ankle-foot orthosis; GR: genu recurvatum.

Table II. Patients’ diagnoses under 3 headings: upper motor neurone pathologies, lower motor neurone pathologies, and traumatic musculoskeletal lesions.

Diagnoses	n (%)
Upper motor neurone pathologies	10 (37)
Stroke	4
Multiple sclerosis	2
Neuro-Behçet’s disease	1
Compressive medullary tumour from T1 to T5	1
Spastic hemiparesis after surgery on an epidermoid cyst on the 4 th ventricular cavity	1
Strumpell-Lorrain disease	1
Lower motor neurone or neuromuscular pathologies	14 (52)
Poliomyelitis	11
Charcot-Marie-Tooth disease	1
Post-radiotherapy plexitis in L3–S1	1
Becker’s disease	1
Traumatic musculoskeletal lesions	3 (11)
Articular destruction after multiple falls	1
Articular destruction after motorcycle accident: posterior knee luxation	1
Gunshot to the femur	1
Total	27 (100)

stance axis (posterior offset KAFOs). In these orthoses, an anterior restriction was used in order to maintain the GR in the range in which the patient had no pain and to prevent falls that were liable to occur if the knee extensors were not strong enough. Six KAFOs had a locked knee joint.

Primary outcomes

Concerning knee pain before and after being fitted with the KAFO, all patients reported an improvement, with the VNRS pain score improving from a median of 85/100 (first quartile = 75; third quartile = 90; range 50–100) to 25/100 (first quartile = 10; third quartile = 35; range 0–75) ($p \leq 0.001$), and the VRS pain assessment improving from “extreme” (median = 4; range 2–4) to “mild” pain (median = 1; range 0–3) ($p \leq 0.001$). These results are shown in Table IV.

Secondary outcomes

The QUEST yielded a total score of 4.0 (SD 0.5), corresponding to “quite satisfied”. The results for each item are shown in Fig. 2.

The median values of the subjectively assessed gait parameters without and with the KAFO were:

- walking distance: “improved” (median = +1; range = 0–2).
- walking speed: “no difference” (median = 0; range = –2–2).
- sensation of recurvatum when walking: “much improved” (median = +2; range = 1–2).
- self-confidence while walking: “improved” (median = +1; range = –2–2).
- feeling safe when walking: “much improved” (median = +2; range = –2–2).
- these results are shown in Table V.

Table III. Patients' range of motion (ROM) and muscular strength

Patients	Recurvatum right/left (°)	ROM hip right/left (°)		ROM ankle right/left (°)		Muscle strength					
		Flexion	Extension	Flexion		Knee extensors MRC	Knee flexors MRC	Knee extensors MRC central	Knee flexors MRC central	MAS Quadriceps /5	MAS Triceps surae /5
1	5	110	15	-5		3	4				
2	30/30	120/120	15/15	20/20		1/2	1/2				
3	30	60	-5	-15		2	2				
4	20	90	-5	0		4	5				
5	20	120	10	0				2	3	2	3
6	25	90	0	5				3	3	1	3
7	20	100	5	5				2	2	2	3
8	10	90	-10	-25				1	2	2	3
9	10	95	-5	5		4	4				
10	10	100	-5	5				3	3	1	3
11	10	110	5	0				2	2	1	3
12	10	90	0	0		2	3				
13	10	120	10	15		5	5				
14	10	100	0	0		3	3				
15	35/35	100/95	10/5	-10/-15		3/2	3/3				
16	30	110	5	-5				3	3	1	2
17	10/5	100/90	10/10	0/5		1/1	1/2				
18	10	100	5	5		3	4				
19	10	110	10	0				3	3	1	2
20	30	120	10	15				2	3	1	2
21	20	90	-10	-10		1	2				
22	30	120	5	-15		2	3				
23	20	100	5	-10		2	3				
24	20	120	10	-10		3	3				
25	20	x	x	x		x	x				
26	15/10	100	0	0		1	0				
27	5	x	x	15				3	3	1	1

Hip and ankle range of motion measured with a goniometer (°).

Muscular strength according MRC testing for lower motor neurone pathologies and traumatic musculoskeletal lesions and upper motor neurone pathologies. (0: no movement, 1: draft voluntary order, 2: voluntary movement included in a synkinetic schema, 3: selective and quantifiable voluntary movement.)

Modified Ashworth Scale (MAS) /5 in triceps surae and quadriceps muscles.

x: missing data; MRC: Medical Research council.

The mean time spent wearing the orthosis per 24 h was 9.8 h (SD 4.4; range: 2–17). Patients reported that they used their orthosis every day from when they got up in the morning until they went to sleep, in their everyday activities. They had to remove the KAFO to wash themselves and sometimes when using the toilet, because of the size of the orthosis or because it damaged their trousers.

Six patients out of 33 did not fit the KAFO: 2 did not answer any phone calls or e-mails and 4 discontinued using the KAFO before 1 month. Three of the patients refused to fit the orthoses due to its weight, discomfort, and difficulty of use, one of them because the KAFO was inducing instability.

The frequency of fallers (yes or no) was 67% before using a KAFO and 37% afterwards ($p=0.021$).

DISCUSSION

The aim of this retrospective study was to analyse the efficiency of KAFOs for alleviating painful GR, and users' satisfaction with this particular choice of treatment. The results show that KAFO is an efficient means of treating posterior knee pain, since it reduced the mean VNRS pain score from 85/100 to 25/100 and the VRS pain assessment from "extreme" to "mild", and the users expressed satisfaction (based on the QUEST), with a mean score of 4.0, corresponding to "quite satisfied", whatever their diagnosis and the cause of the GR. The subjective gait parameters and the patients' impression of GR also improved. Based on the QUEST, the KAFOs rated well in terms of safety, durability and effectiveness, whereas their comfort, weight and

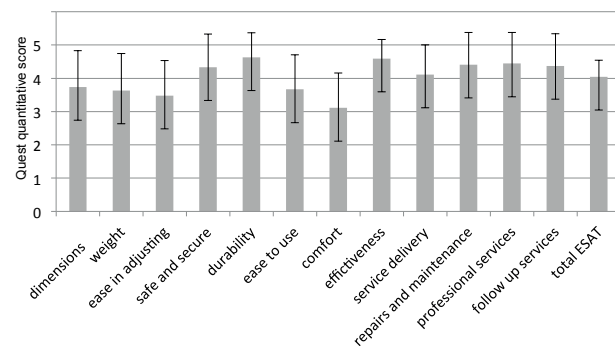


Fig. 2. Results of the Quebec User Evaluation of Satisfaction with assistive Technology (QUEST 2.0). Mean scores obtained on each of the items in the French version of QUEST 2.0 with standard deviations. (QUEST quantitative score: 1: "not satisfied at all"; 2: "not very satisfied"; 3: "more or less satisfied"; 4: "quite satisfied"; 5: "very satisfied").

Table IV. Main outcomes: patients' mean knee pain assessments before and after wearing a knee-ankle-foot orthosis

Pain assessment	Before using KAFO	After using KAFO	p-value
VNRS, medians (range)	85/100 (50–100)	25/100 (0–75)	≤ 0.001
VNRS, medians	4, "extreme" pain	1, "mild" pain	≤ 0.001
	4: 15/27	4: 0/27	
	3: 10/27	3: 1/27	
	2: 2/27	2: 6/27	
	1: 0/27	1: 13/27	
	0: 0/27	0: 7/27	

Results of pain assessments made in the present study with: the verbal numerical rating scale (VNRS) and verbal rating scale (VNRS) before and after using a knee-ankle-foot orthosis (KAFO). ("no pain" [0], "mild pain" [1], "moderate pain" [2], "severe pain" [3] and "extreme pain" [4]).

Table V. Median values of the subjectively assessed gait parameters* without and with the knee-ankle-foot orthosis (KAFO)

Gait parameters	
Walking distance	1 (improved)
Walking speed	0 (no difference)
Sensation of recurvatum	2 (much improved)
Self-confidence when walking	1 (improved)
Safety when walking	2 (much improved)

*Subjective gait parameters were rated without and with the KAFO as: "much improved" (+2), "improved" (+1), "no difference" (0), "worse" (-1), "much worse" (-2) (15, 19), and only scored by patients.

dimensions were given the lowest ratings, confirming the clinical assessment and users' criticisms.

The patients reported that they used their KAFOs every day, removing them only for sleeping, bathing, and getting dressed, and that they wore them for up to 9 h per day, which is quite long. In 2 patients with a recurvatum of only 5°, the VNRS knee pain score decreased from 85 and 80/100 to 60 and 20/100, respectively, which shows that the intensity of the pain does not necessarily depend on the severity of the GR. It should be noted that included patients had worn the orthosis for more than 3 months since the last correction. The latter criterion was adopted in order to study the effects of the final orthoses, but it also shows that when orthoses are accepted, it is with efficiency.

The study population (27 patients) was rather small, which was a limitation of this study, but all previous studies on orthotic devices for patients with GR have been conducted on rather small populations (between 11 and 25 patients (1, 14, 20, 22)) because it is difficult to find larger populations of patients with an indication of KAFO for painful GR. Moreover, the strict inclusion criteria used here resulted in the exclusion of all patients with instability, in order to assess the effects of KAFOs on pain and not on instability, although both symptoms often occur concurrently. This is another reason for the small sample of patients in this study. A further limitation of the present study is that, since patients with musculoskeletal and neurological diseases were included, a heterogeneous population was obtained, which is typical of KAFO users. This heterogeneity confirmed the sig-

nificance of our results. However, for an homogenous group, such as the group of patients with poliomyelitis, it was observed that wearing KAFOs reduced their knee pain scores from 83/100 on the VNRS scale to 29/100, and good ratings of patients' satisfaction (QUEST=4.1) were obtained, even on the item "comfort" (3.5/5). Although poliomyelitis patients in general do not have sensory disorders, they were highly sensitive to the pressure exerted by the orthosis (38, 39).

This retrospective study is the first step in the second part of a larger ongoing study with a prospective design. The retrospective approach included assessment after the KAFO was fitted, which may influence the results. Patients included in this study were fitted with a KAFO because of the efficiency of the device. In the second part of the study, using a prospective approach, data will be collected before making the decision to apply orthoses, and kinematic parameters (based on optoelectronic technology with a Helen Hayes marker set up (26)) will be recorded and used to define the cause of the GR and calculate the mechanical energy expenditure (40) and the efficiency of the angular correction before and after orthotic correction. The second question to address in order to improve the medical management of GR, is what range of motion of the knee is required with the KAFO in order to reduce knee pain but not create instability of the knee.

Conclusion

Treating painful posterior GR with KAFOs was found to be an efficient means of alleviating pain, regardless of the patients' diagnoses. This method met with a good level of patient satisfaction. Further research is needed in order to develop an effective approach to management of GR.

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REFERENCES

- Appasamy M, De Witt ME, Patel N, Yeh N, Bloom O, Oreste A. Treatment strategies for genu recurvatum in adult patients with hemiparesis: a case series. *PM&R* 2015; 7: 105–112.
- Perry J. Lower-extremity bracing in hemiplegia. *Clin Orthop* 1969; 63: 32–38.
- Gross R, Delporte L, Arsenault L, Revol P, Lefevre M, Cleve-not D, et al. Does the rectus femoris nerve block improve knee recurvatum in adult stroke patients? A kinematic and electromyographic study. *Gait Posture* 2014; 39: 761–766.
- Cooper A, Alghamdi GA, Alghamdi MA, Altowaijri A, Richardson S. The relationship of lower limb muscle strength and knee joint hyperextension during the stance phase

- of gait in hemiparetic stroke patients. *Physiother Res Int* 2012; 17: 150–156.
5. Fish DJ, Kosta CS. Genu recurvatum: identification of three distinct mechanical profiles. *JPO J Prosthet Orthot* 1998; 10: 26–32.
 6. Morris ME, Bach TM, Goldie PA. Electrogoniometric feedback: its effect on genu recurvatum in stroke. *Arch Phys Med Rehabil* 1992; 73: 1.
 7. Loudon JK, Goist HL, Loudon KL. Genu recurvatum syndrome. *J Orthop Sports Phys Ther* 1998; 27: 361–367.
 8. Hogue R, McCandless S. Genu recurvatum: auditory biofeedback treatment for adult patients with stroke or head injuries. *Arch Phys Med Rehabil* 1983; 64: 368–370.
 9. Stanic U, Acimović-Janežic R, Gros N, Trnkoczy A, Bajd T, Kljajić M. Multichannel electrical stimulation for correction of hemiplegic gait. Methodology and preliminary results. *Scand J Rehabil Med* 1978; 10: 75.
 10. Novak AC, Olney SJ, Bagg S, Brouwer B. Gait changes following botulinum toxin A treatment in stroke. *Top Stroke Rehabil* 2009; 16: 367–376.
 11. Pradon D, Hutin E, Khadir S, Taiar R, Genet F, Roche N. A pilot study to investigate the combined use of Botulinum toxin type-a and ankle foot orthosis for the treatment of spastic foot in chronic hemiplegic patients. *Clin Biomech* 2011; 26: 867–872.
 12. Fatone S, Gard SA, Malas BS. Effect of ankle-foot orthosis alignment and foot-plate length on the gait of adults with poststroke hemiplegia. *Arch Phys Med Rehabil* 2009; 90: 810–818.
 13. Esquenazi A, Ofluoglu D, Hirai B, Kim S. The effect of an ankle-foot orthosis on temporal spatial parameters and asymmetry of gait in hemiparetic patients. *PM&R* 2009; 1: 1014–1018.
 14. Farncombe P. The Swedish knee cage. Management of the hyperextended hemiplegic knee. *Physiotherapy* 1980; 66: 33.
 15. Portnoy S, Frechtel A, Raveh E, Schwartz I. Prevention of genu recurvatum in poststroke patients using a hinged soft knee orthosis. *PM&R* 2015; 7: 1042–1051.
 16. Pinzur M, Sherman R, DiMonte-Levine P, Kett N, Trimble J. Adult-onset hemiplegia: changes in gait after muscle-balancing procedures to correct the equinus deformity. *J Bone Joint Surg Am* 1986; 68: 1249–1257.
 17. Caillet F, Mertens P, Rabaseda S, Boisson D. The development of gait in the hemiplegic patient after selective tibial neurotomy. *Neurochirurgie* 1998; 44: 183–191.
 18. Bleyenheuft C, Bleyenheuft Y, Hanson P, Deltombe T. Treatment of genu recurvatum in hemiparetic adult patients: a systematic literature review. *Ann Phys Rehabil Med* 2010; 53: 189–199.
 19. Bettoni E, Ferriero G, Bakhsh H, Bravini E, Massazza G, Franchignoni F. A systematic review of questionnaires to assess patient satisfaction with limb orthoses. *Prosthet Orthot Int* 2016; 40: 158–169.
 20. Morinaka Y, Matsuo Y, Nojima M, Morinaka S. Clinical evaluation of a knee-ankle-foot-orthosis for hemiplegic patients. *Prosthet Orthot Int* 1982; 6: 111–115.
 21. Morinaka Y, Matsuo Y, Nojima M, Inami Y, Nojima K. Biomechanical study of a knee-ankle-foot-orthosis for hemiplegic patients. *Prosthet Orthot Int* 1984; 8: 97–99.
 22. Boudarham J, Zory R, Genet F, Vigné G, Bensmail D, Roche N, et al. Effects of a knee-ankle-foot orthosis on gait biomechanical characteristics of paretic and non-paretic limbs in hemiplegic patients with genu recurvatum. *Clin Biomech* 2013; 28: 73–78.
 23. Kerrigan DC, Deming LC, Holden MK. Knee recurvatum in gait: a study of associated knee biomechanics. *Arch Phys Med Rehabil* 1996; 77: 645–650.
 24. Simon S, Deutsch S, Nuzzo R, Mansour M, Jackson J, Koskinen M, et al. Genu recurvatum in spastic cerebral palsy. Report on findings by gait analysis. *J Bone Joint Surg Am* 1978; 60: 882–894.
 25. Olney SJ, Monga TN, Costigan PA. Mechanical energy of walking of stroke patients. *Arch Phys Med Rehabil* 1986; 67: 92–98.
 26. Zissimopoulos A, Fatone S, Gard SA. Biomechanical and energetic effects of a stance-control orthotic knee joint. *J Rehabil Res Dev* 2007; 44: 503.
 27. Merkel K, Miller N, Westbrook P, Merritt J. Energy expenditure of paraplegic patients standing and walking with two knee-ankle-foot orthoses. *Arch Phys Med Rehabil* 1984; 65: 121–124.
 28. Hachisuka K, Makino K, Wada F, Saeki S, Yoshimoto N. Oxygen consumption, oxygen cost and physiological cost index in polio survivors: a comparison of walking without orthosis, with an ordinary or a carbon-fibre reinforced plastic knee-ankle-foot orthosis. *J Rehabil Med* 2007; 39: 646–650.
 29. Astephen JL, Deluzio KJ, Caldwell GE, Dunbar MJ, Hubley-Kozey CL. Gait and neuromuscular pattern changes are associated with differences in knee osteoarthritis severity levels. *J Biomech* 2008; 41: 868–876.
 30. Andriacchi TP, Mündermann A, Smith RL, Alexander EJ, Dyrby CO, Koo S. A framework for the in vivo pathomechanics of osteoarthritis at the knee. *Ann Biomed Eng* 2004; 32: 447–457.
 31. Andriacchi TP, Koo S, Scanlan SF. Gait mechanics influence healthy cartilage morphology and osteoarthritis of the knee. *J Bone Joint Surg* 2009; 91: 95–101.
 32. Arazpour M, Bani MA, Hutchins SW. Reciprocal gait orthoses and powered gait orthoses for walking by spinal cord injury patients. *Prosthet Orthot Int* 2013; 37: 14–21.
 33. Hjermstad MJ, Fayers PM, Haugen DF, Caraceni A, Hanks GW, Loge JH, et al. Studies comparing numerical rating scales, verbal rating scales, and visual analogue scales for assessment of pain intensity in adults: a systematic literature review. *J Pain Symptom Manage* 2011; 41: 1073–1093.
 34. Demers L, Monette M, Lapierre Y, Arnold D, Wolfson C. Reliability, validity, and applicability of the Quebec User Evaluation of Satisfaction with assistive Technology (QUEST 2.0) for adults with multiple sclerosis. *Disabil Rehabil* 2002; 24: 21–30.
 35. Eckhardt MM, Mulder MCB, Horemans HL, van der Woude LH, Ribbers GM. The effects of high custom made shoes on gait characteristics and patient satisfaction in hemiplegic gait. *Gait Posture* 2011; 34: 543–547.
 36. Tyson S, Thornton H. The effect of a hinged ankle foot orthosis on hemiplegic gait: objective measures and users' opinions. *Clin Rehabil* 2001; 15: 53–58.
 37. Demers L, Weiss-Lambrou R, Ska B. Item analysis of the Quebec user evaluation of satisfaction with assistive technology (QUEST). *Assist Technol* 2000; 12: 96–105.
 38. Waring WP, Maynard F, Grady W, Grady R, Boyles C. Influence of appropriate lower extremity orthotic management on ambulation, pain, and fatigue in a postpolio population. *Arch Phys Med Rehabil* 1989; 70: 371–375.
 39. Genêt F, Schnitzler A, Mathieu S, Autret K, Théfenne L, Dizien O, et al. Orthotic devices and gait in polio patients. *Ann Phys Rehabil Med* 2010; 53: 51–59.
 40. Caty GD, Detrembleur C, Bleyenheuft C, Lejeune TM. Reliability of lower limb kinematics, mechanics and energetics during gait in patients after stroke. *J Rehabil Med* 2009; 41: 588–590.