Inertial sensors as measurement tools of elbow range of motion in gerontology

Background and purpose: Musculoskeletal system deterioration among the aging is a major reason for loss of autonomy and directly affects the quality of life of the elderly. Articular evaluation is part of physiotherapeutic assessment and helps in establishing a precise diagnosis and deciding appropriate therapy. Reference instruments are valid but not easy to use for some joints. The main goal of our study was to determine reliability and intertester reproducibility of the MP-BV, an inertial sensor (the MotionPod® [MP]) combined with specific software (BioVal [BV]), for elbow passive range-of-motion measurements in geriatrics.

Methods: This open, monocentric, randomized study compared inertial sensor to inclinometer in patients hospitalized in an acute, post-acute, and long-term-care gerontology unit.

Results: Seventy-seven patients (mean age 83.5±6.4 years, sex ratio 1.08 [male/female]) were analyzed. The MP-BV was reliable for each of the three measurements (flexion, pronation, and supination) for 24.3% (CI 95% 13.9–32.8) of the patients. Separately, the percentages of reliable measures were 59.7% (49.2–70.5) for flexion, 68.8% (58.4–79.5) for pronation, and 62.3% (51.2–73.1) for supination. The intraclass correlation coefficients were 0.15 (0.07–0.73), 0.46 (0.27–0.98), and 0.50 (0.31–0.98) for flexion, pronation, and supination, respectively.

Conclusion: This study shows the convenience of the MP-BV in terms of ease of use and of export of measured data. However, this instrument seems less reliable and valuable compared to the reference instruments used to measure elbow range of motion in gerontology.

Keywords: reliability, intertester reproducibility, inclinometer

Introduction

Musculoskeletal system deterioration among the aging is a major reason for loss of autonomy and directly affects the quality of life of the elderly.1–3 In this context, physiotherapeutic assessment helps in evaluating elderly patient deficiencies and thus in planning an appropriate rehabilitation program. Articular evaluation is part of this assessment and allows joint deviation quantification, identification of orthopedic deformities, observation of painful events, and follow-up of progress made by patients. It also helps in establishing a precise diagnosis and deciding appropriate therapy.4

An essential stage of this evaluation is the measurement of joint angles. Within a clinical setting, physiotherapists do have technical references: the goniometer and inclinometer are valid instruments;5–7 however, positioning of these tools is not easy with some joints.8–10 Other techniques are also used in tests: the meter ribbon (easy use for estimating improvement in range of motion [ROM] but not adapted to a unique assessment) and visual assessment (often practiced in clinical tests but subjective and not entirely replicable).11,12 It seemed necessary to find an instrument that would allow a combination of all the assets of those reference techniques—that is, an instrument that is reliable and replicable but also simple, fast, and easy to use.
Actigraphy is a technique for measuring locomotive activity. Traditionally, actimeters were composed of single piezoelectric accelerometers worn on the waist. With the improvement of technology in the past 30 years, actimeters have become more precise (with multiple piezoelectric accelerometers) and the position of wear has been diversified (wrist, chest, hip) in response to new applications. Several practical applications of actigraphy have been proposed, such as assessment of hyperactivity in children and behavioral disorders among patients suffering from dementia; ecological assessment of daily living activities; assessment of sedative effects of some drugs; and screening for certain sleep problems. However, the actigraph can also be used for measurement of articular ROM. Indeed, the actigraph, in combination with specific software, is used as an inertial sensor for measurement of lower limb articulations but also for cervical spine.

The main goal of the present study was to determine reliability, with an inclinometer as reference, and intertester reproducibility of an actigraph that uses an inertial sensor (the MotionPod; Movea SA, Grenoble, France) associated with specific software (BioVal; RM Ingénierie, Groupe Cegedim, Rodez, France) for obtaining elbow passive ROM measurements in geriatrics. In this study, we also evaluated the duration of examination and the acceptability of the instrument according to the patient and the physiotherapist. From this point onward, we shall use the term “MP-BV” to define the actigraph/software coupling.

Methods
Location and subjects
This open, monocentric, randomized study (NCT01462760) took place from October 1, 2011 to May 1, 2012. Patients were recruited within the acute, post-acute, and long-term care gerontology unit of University Hospital of Nice (Nice, France). The study was approved by the South Mediterranean V Ethical Review Board (Comité de Protection des Personnes Sud Méditerranée V), as well as by the French Agency for the Safety of Health Products (ANSM). Patients who agreed to participate gave their written informed consent.

Included patients were 65 years or older and had received a physiotherapy prescription from the physiotherapists of Department of Geriatrics mentioning the need for a passive joint assessment. Patients who agreed to participate had to be covered by social insurance. Exclusion criteria were: patients suffering from a pathology for which heating of joints was contraindicated; patients wearing a pacemaker, a metallic prosthesis, or osteosynthesis equipment located less than 10 cm from the captor; patients incapable of maintaining a straight wrist position and/or a neutral shoulder position; and patients having a joint deviation in flexion, extension, pronation or supination of less than 20°.

Recruitment and randomization
Patients were pre-recruited during weekly physiotherapy sessions at the Department of Geriatrics. Due to organizational constraints, a maximum of six patients per week were included. An explanation of the study and an associated pamphlet were given to the patients by the physician responsible for the study, with a 7-day retraction delay. The inclusion visit began with signing of consent and was led by the physician after those 7 days.

At the end of the inclusion visit, an inclusion form was completed by the physician. This form included the first and last name of the patient and the confirmation of all inclusion and non-inclusion criteria. Then, this form was forwarded to a clinical research assistant of the Clinical Research and Innovation Department (DRCI), who randomly assigned the patients. The randomization was centralized at the methodological center and done in a 1:1 ratio. The generation of the allocation sequence was made with number permutation tables in block sizes of six. Consequently, among six patients, three were allocated to room A and three were allocated to room B, and so on. All patients were examined successively in both rooms. Furthermore, in room A, the running order on both instruments (MP-BV and inclinometer) was also randomized. Physiotherapists were also randomly assigned to room A or B (Figure 1).

Materials used for the measures
The MotionPod is a patented hardware solution for motion sensing that features a state-of-the-art miniaturized motion-sensing microelectromechanical system. It contains three-axis accelerometers, three-axis magnetometers, and three-axis gyroscopes in a compelling form factor the size of a standard wristwatch (33×21×15 mm [1.3×0.8×0.6”]) and weighs 14 g (0.5 oz). The battery was a 150 mAh Li-ion polymer technology battery with a battery life of 6 hours and a charging duration of 2–3 hours (500 charging cycles). The device is worn by a patient by way of a bracelet attached to the mechanical interface. The sampling rate for accelerometers, gyroscopes, and magnetometers was adjustable between 25 and 200 Hz with 12-bit resolution. The information from the MotionPod was transferred via radio waves to the MotionPod controller, which was linked to a computer with a USB cable. The wireless band ranged from 10 m (30 feet) to 30 m (100 feet), depending on the environment, with a frequency band of 2.4 GHz. The measurement range was a full 360°. The fusion of data from accelerometers, magnetometers, and gyroscopes was done with an application programming.
An inertial sensor for measuring elbow range of motion

The inclinometer used as reference was the Base- 
line® Bubble inclinometer (Fabrication Enterprises, 
With Plains, NY, USA), the dimensions of which were 
101.6×5.08×101.6 mm (4×0.2×4″) for 90.72 g (0.2 lbs). For 
the measure, the inclinometer was placed near the elbow. The 
physiotherapist turned the dial until the scale read 0, then took 
the elbow through its range of motion. Finally, the range was 
read directly from the dial.

**Experimentation process**

Experimentation was performed in the Physiotherapy Unit 
of the Gerontology Department over one half-day per week. 
A training session on both instruments used in this study was 
held by physiotherapists before the study began so that all 
four investigators had the same level of knowledge regarding 
the use of the MP-BV. No physiotherapist had had any expe-
rience with any inertial sensor before the training sessions.

Measurements were made on the elbow joint, as it allows 
good validity and intertester reproducibility.1 Physiothera-
pists successively measured flexion, pronation, and supina-
tion. In room A, experimentation included measurement 
with the MP-BV as well as with the inclinometer. The screen 
displaying the result was hidden from the physiotherapist so 
as not to influence the inclinometer measurement. In room B, 
the experiment included a measurement with the MP-BV, in 
order to assess intertester reliability. Considering that incli-
nometer was the instrument of reference, only one measure 
was taken. For MP-BV two measures were taken for each 
patient to assess intertester reliability, which were performed 
by different physiotherapists for each measure.

When the passive elbow flexion was measured, the arm 
was placed vertically and the forearm was placed in supi-
nation in the same straight line as that formed by the arm. 
The inclinometer was placed on the longitudinal axis of the 
mobile segment, on the dorsal surface, and on the middle of 
the forearm. For this measurement, we used two MotionPod 
devices; the first one, fixed on the wrist, measured the joint 
angle, and the second one, on the arm, was used as reference 
positioning (Figure 2).

Regarding pronosupination, the arm was placed 
vertically; the forearm was placed horizontally with the palm 
of the hand up, the thumb upward, and the longitudinal axis 
of the third metacarpus in the same axis as the forearm. The 
inclinometer was positioned on the dorsal face of the wrist 
and the MotionPod the motion pod was positioned on the 
external face of the wrist.
Variables studied
Measurement reliability
Passive elbow ROM measurements expressed in degrees and obtained with the MP-BV were compared to those obtained with an inclinometer for all three of the movements (flexion, pronation, and supination). We excluded elbow extension because the articular deviation of this joint is inferior to 10°, which is inferior to the variability accepted between the two techniques. A 10° scope of error is acceptable for goniometric measurements. Consequently, we considered as reliable a measure with the MP-BV with a difference of less than 10° of the measure obtained with the inclinometer. The percentage of reliable measure was calculated in this way for each movement.

Intertester reproducibility
Intertester reproducibility was assessed through two series of passive ROM measurements conducted with the MP-BV by two researchers. For each of the three movements, intertester reproducibility of MP-BV measurement was evaluated by calculating the intraclass correlation coefficient (ICC). Reproducibility was also evaluated through the rates of concordant measures with both techniques (less than 10° difference between the measures) for each movement.

Duration of examination
The durations of examination (in seconds) obtained with both instruments were compared. Duration was defined as the time at which the patient was ready for examination until the time at which the data were recorded in the patient’s file.

Acceptability
The patient rated the comfort of the measurement process for both instruments using a ten-point visual analog scale (VAS). Meanwhile, each physiotherapist rated his acceptance of the instruments using a 10-point VAS, as well as the ease of use and of data exporting of both instruments.

Statistical analysis
It was decided a priori that MP-BV would be considered a reliable measurement technique if the three joint angles from at least 90% of patients had less than 10° error when compared to the inclinometer. Considering the assumption that 95% of patients would have a valid measure, it was necessary to include 73 patients so that the lower limit of the 95% confidence interval (CI) would be above 90%. To account for potential measurement failures, the decision was made to include 84 patients.

First, a descriptive analysis was made by the biostatistician from the DRCl of Nice University Hospital. Results were presented as mean ± standard deviation for quantitative variables and percentage for qualitative variables. The percentage of patients with a valid global assessment was calculated and presented with the corresponding 95% CI as well as percentages of reliable measurements for each movement (flexion, pronation, supination). An ICC value <0.40 was considered poor, 0.40–0.50 moderate, 0.50–0.70 good, and 0.70–1.0 excellent.

Examination duration was compared between the two techniques using the Student’s t-test for paired data. For this analysis, only data recorded in room A were used, as examinations were performed there with both techniques and by the same physiotherapist.

Finally, for acceptability analyses, VAS scores (recorded in room A) were compared using the Student’s t-test for paired data. All the tests were two-sided and the significance level was set to 5%. Statistical analyses were performed using SAS Enterprise Guide (v 4.1; SAS Institute Inc., Cary, NC, USA).

Results
Three hundred and fifteen patients were screened during weekly physiotherapy sessions. Of these, 84 patients were deemed eligible for inclusion in the study, and 77 of these 84 (91.7%) were eventually included in the validation study (Figure 3).

Characteristics of the patients with missing data for analyses did not differ significantly from analyzable patients. For these 77 patients, the mean age was 83.5±6.4 (range 68–101) years; 40 (52.0%) were women and 37 (48.0%) men, resulting in a sex ratio of 1.08.
Measuring reliability
The MP-BV was reliable on each of the three measurements (flexion, pronation, and supination) for 24.3% (13.9–32.8) of the patients. Separately, the percentages of reliable measures were 59.7% (49.2–70.5) for flexion, 68.8% (58.4–79.5) for pronation, and 62.3% (51.2–73.1) for supination (Table 1).

Intertester reproducibility
The ICCs were 0.15 (0.07–0.73) for flexion, 0.46 (0.27–0.98) for pronation, and 0.50 (0.31–0.98) for supination. For the same movements, respectively, the concordant measures percentages were 53.2% (42.1–64.4), 57.1% (46.1–68.2), and 53.2% (42.1–64.4).

Duration of examination
Mean examination duration was significantly longer with the MP-BV than with the inclinometer, at 117.4 (±26.9) seconds and 104.4 (±34.0) seconds, respectively (P<0.0001).

Acceptability
There was no significant difference between the two instruments regarding patient comfort, with a mean VAS score for the MP-BV of 9.21 (±1.10) versus 9.26 (±0.99) for the inclinometer (P=0.49, DF=76, t=0.69). Results were the same regarding the physiotherapist’s acceptance of the instruments, with a mean VAS score of 7.88 (±1.97) for the MP-BV and 7.70 (±1.70) for the inclinometer (P>0.05, DF=76, t=1.43).

Adverse events
No adverse events were recorded during the study.

Discussion
Measuring articular ROM requires the use of various instruments. However, even those instruments provide reliable and replicable measures, they present difficulties in the use and the use of reference instruments (like inclinometer or goniometer) required that the user was very rigorous in positioning the instrument before using it.10 The purpose of this study was...
to determine the reliability and the intertester reproducibility of the MP-BV as a new measurement instrument for ROM, which may be simpler to use than other instruments.

The results of this study show the convenience of the MP-BV in terms of ease of use and of export of measured data. However, this instrument seems less reliable and useful compared to the reference instruments used to measure elbow ROM in gerontology.

The choice of measurement instrument in clinical practice is not consensual, and study results differ between authors. For this reason, van de Pol et al. concluded in their literature review that more research regarding these instruments was necessary. In our study, we chose the inclinometer as a reference instrument due to its validity, but also because of its higher intertester reproducibility compared to a classical goniometer on the elbow joint. Moreover, the inclinometer is a fast and easy-to-use instrument with which physiotherapists from the Nice University Hospital are familiar. Finally, an important physiological valgus could hamper common goniometer implementation on the elbow joint. Chapleau et al. highlighted in their study the need for using radiological measurement in clinical protocols. However, radiological measurement was difficult to apply in our study.

Remarkably, this study analyzed 77 patients, which represents a significant number of patients compared to similar studies which have validated reference instruments. Within the context of the study, a double-blind experiment for measurements was not achievable.

We experienced some difficulties in measurement that could explain the differences between the two instruments’ outcomes. Elbow flexion measurement is usually done by placing the shoulder in a neutral position, and the choice of using two inertial sensors was made to avoid subject compensation while performing this measure (shoulder protraction). Unlike the inclinometer, the MP-BV was attached to the skin; however, skin elasticity decreases with age and, as a result, there was a low adjustment of the sensor in regard to the underlying bony landmark, creating a measurement difference higher than 10°. This particularity could also explain measurement differences when pronation and supination movements were performed.

Contrary to what was expected, the importance of a rigorous marked of reference point proved to be extremely important. Indeed, the positioning of the sensor on the joint by both physiotherapists was not exactly the same and may have been responsible for the different outcomes. Moreover, a clamping system was not provided with the MotionPod, and it was necessary to use a skin guard to protect the elderly patients’ skin, which is often fragile, increasing tracking difficulties.

Regarding examination duration, it is important to note that this not only included examination duration, but also the setting-up of the skin guard, thus increasing the total measuring time with the MP-BV compared to the inclinometer.

Conclusion

With the continuing development of new technologies, we shall witness more and more new instruments that will require assessment using similar protocols to those used for medications. Based on this study, the MP-BV seems less reliable and to give less reproducible measures than one of the reference instruments used to assess elbow ROM in the elderly. Taking into account its limitations as well as its ease of both use and data exportation, it could be interesting to compare MP-BV measurements to radiographic measurements in a younger population.

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Author contributions

All authors contributed toward data analysis, drafting and revising the paper and agree to be accountable for all aspects of the work.

Disclosure

This study benefited from Ministry of Industry funding within the Centre for Innovation and Health Usage (Centre d’Innovation et d’Usage en Santé [CIU-S]). MP-BV developers were part of the initial CIU-S project consortium. The statistical outcomes analysis was independently realized by the DRCI and the data interpretation was performed by physicians from the public hospital. Participants for this study (physicians, physiotherapists) who were involved in this research protocol and in the interpretation of the outcomes did not receive any specific compensation. The authors report no other conflicts of interest in this work.
References


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