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Rater Agreement on Gait Assessment during Neurologic Examination of Horses

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Background: Reproducible and accurate recognition of presence and severity of ataxia in horses with neurologic disease is important when establishing a diagnosis, assessing response to treatment, and making recommendations that might influence rider safety or a decision for euthanasia.

Objectives: To determine the reproducibility and validity of the gait assessment component in the neurologic examination of horses.

Animals: Twenty-five horses referred to the Royal Veterinary College Equine Referral Hospital for neurological assessment (n = 15), purchased (without a history of gait abnormalities) for an unrelated study (n = 5), or donated because of perceived ataxia (n = 5).

Methods: Utilizing a prospective study design; a group of board-certified medicine (n = 2) and surgery (n = 2) clinicians and residents (n = 2) assessed components of the equine neurologic examination (live and video recorded) and assigned individual and overall neurologic gait deficit grades (0–4). Inter-rater agreement and assessment-reassessment reliability were quantified using intraclass correlation coefficients (ICC).

Results: The ICCs of the selected components of the neurologic examination ranged from 0 to 0.69. “Backing up” and “recognition of mistakes over obstacle” were the only components with an ICC > 0.6. Assessment-reassessment agreement was poor to fair. The agreement on gait grading was good overall (ICC = 0.74), but poor for grades ≤ 1 (ICC = 0.08) and fair for ataxia grades ≥ 2 (ICC = 0.43). Clinicians with prior knowledge of a possible gait abnormality were more likely to assign a grade higher than the median grade.

Conclusion and Clinical Importance: Clinicians should be aware of poor agreement even between skilled observers of equine gait abnormalities, especially when the clinical signs are subtle.

Key words: Agreement; Ataxia; Physical examination; Reliability.

A thorough physical examination in combination with the history is the primary source of initial information for any clinician and influences decision making for further diagnostic and therapeutic intervention. Examinations should therefore have good reliability (how well can patients be distinguished from each other), high agreement (low measurement error for repeated assessments) and results of the examination should be valid (accurate). Inaccurate or unreliable assessment of an underlying problem might lead to misdiagnosis or inappropriate further testing and treatments. Consequently, inherent limitations of the physical examination because of its subjectivity should be recognized and the examination optimized to reduce variability and bias.

Few studies have examined reproducibility and validity of physical examinations in horses and humans. There is expectation bias in gait and lame-ness assessment of horses despite good agreement between observers where a lameness score (AAEP, 0–5) was >1.5/5 (k = 0.86), or low agreement (k = 0.23), for lameness grades ≤1.5/5. Low agreement is especially of concern when additional testing options are limited and when decisions are made relevant to human safety or animal euthanasia. One such example is neurologic examination of horses where...

Abbreviations:
- CMID: clinically minimal important difference
- CVM: cervical vertebral malformation-malarticulation
- ERH: Equine Referral Hospital
- ICC: intraclass correlation coefficient
- PCA: principal component analysis
- RVC: The Royal Veterinary College
- SARA: scale for the assessment and rating of ataxia
diagnostic imaging and other testing options are limited by physical constraints, by poor inherent sensitivity or specificity, or both and where the prognosis for recovery often is poor.

During assessment of gait in neurologic examination of horses, clinicians typically grade the severity of the neurologic deficit according to a modified grading scale (0–5), where grade 0 is assigned to horses without neurologic deficits and grade 5 is assigned to horses that are recumbent. Despite general acceptance and application of this system, its reproducibility and validity have never been tested; neither has there been any formal assessment of the role for expectation bias in neurologic examination of horses.

The aim of this study was to evaluate the subjective assessment of horses with and without varying neurologic deficits by evaluating the rater and assessment-reassessment agreement of a modified ataxia-grading scale. In addition, we aimed to determine whether expectation bias plays a role in neurologic assessment of horses. We hypothesized that (1) agreement between raters (inter-rater) is good when assessing each part of the neurologic examination of horses with and without suspected neurologic deficits; (2) assessment-reassessment agreement is good when raters assess horses with and without suspected neurologic deficits twice on video; (3) the modified ataxia-grading scale has good agreement applied to horses with moderate to severe ataxia (ataxia grade ≥ 2), but poor agreement for normal horses or horses with low-grade deficits (ataxia grade ≤ 1); (4) raters who know a horse is suspected of having an abnormal gait (unrelated to lameness) assign a higher grade than the median ataxia grade.

Materials and Methods

The project was designed and conducted as a prospective cross-sectional reproducibility study according to the Guidelines for Reporting Reliability and Agreement Studies. The study was approved by the Ethics and Welfare Committee of the Royal Veterinary College and where appropriate, was conducted under specific Home Office License according to the Animal (Scientific Procedures) Act (1986) of the United Kingdom. Please see Data S1 for the full details on methodology.

Raters

Six raters were recruited among clinicians at The Equine Referral Hospital (ERH) at the Royal Veterinary College (RVC), UK: 2 were board-certified internists (Large Animal; ACVIM) and 2 were board-certified surgeons (ECVS or ACVS). Two raters were second-year residents in either large animal internal medicine or equine surgery. Anonymity in scoring was maintained by assigning each rater a random, single digit number unknown to the author responsible for data entry and analysis.

Raters were not given any information about the horse’s signalment, history, presentation or other clinical or clinicopathologic findings before the assessment of gait; however if a clinician had knowledge about the examined horse the rater was asked to disclose knowledge of the history or source of the horse and its reason for presentation.

Horses

Examinations were conducted between October 2010 and November 2012 at RVC ERH. Horses were recruited from 3 sources: Group 1 included horses with no known history of gait abnormalities that were purchased for an unrelated study of recurrent laryngeal neuropathy. Group 2 comprised horses referred to the ERH from first opinion practice for evaluation of gait deficits or perceived ataxia. Horses were recruited to Group 3 if a decision for euthanasia had been made in first opinion practice because of perceived moderate to severe ataxia. Horses that were considered on ethical grounds to be too ataxic to travel were excluded from Group 3. The horses were examined in order of presentation to the ERH.

Neurologic Examination

Every horse underwent a full and identical neurologic gait assessment that included walk and trot in a straight line, walking with the head elevated, walking with a blindfold, and standing tail pull, walking in small circles in both directions, backing up, lateral placement of distal thoracic limbs, crossing over of distal thoracic limbs, hopping on each thoracic limb, bilateral cervicofacial and panniculus reflexes, walking over an obstacle (10 × 20 cm pole), walking up and down a slope with and without head elevation. Raters completed a questionnaire for each step of the examination and were asked if the result was normal or abnormal as well as further characteristics (Table 2 and Data S2). The horses’ gait were graded according to a 5-point ataxia scale first proposed by Mayhew et al and later modified by Reed (Table 1). In a separate question, the horses were assessed and graded for lameness according to an 11-point scale. The grades were assigned after walk and trot in a straight line and rescored with an overall grade, after the complete examination.

Video

All examinations were filmed with a video camera (1080 p, 50 Hz, shutter speed: 1/250 second) with standardized recording and editing (further details in Data S1). A set of 10 videos (median grades: grade 0 [n = 3], grade 1 [n = 2], grade 2 [n = 2], grade 3 [n = 2], and grade 4 [n = 1]) were selected for test-retest reliability and edited. The assessment-reassessment study was based on scoring of these 10 videos on 2 separate occasions, making it

Table 1. The modified ataxia-grading scale used in this study. The text explaining each grade was printed on the questionnaire. No recumbent horses (grade 5) were included in the study.

| Grade | Description |
|-------|-------------|
| 0     | No gait deficits at the walk |
| 1     | No gait deficits identified at the walk and deficits only identified during further testing |
| 2     | Deficits noted at the walk |
| 3     | Marked deficits noted at the walk |
| 4     | Severe deficits noted at the walk and may fall or nearly fall at normal gaits |
a video-to-video comparison to reduce the impact of bias between live and video assessment.

**Postmortem Examination and Histopathology**

For all euthanized horses, the entire spinal cord was removed and fixed. Sections of the spinal cord were examined segmentally at the level of the dorsal root from C1-T2 with additional sections examined at T9, T16, and L5. A Diplomate of the European College of Veterinary Pathology with a specific interest in neuropathology (author KB) examined the transverse and longitudinal sections that were processed and stained with hematoxylin and eosin and Luxol fast blue. The pathologist was blinded to the signalment, history, clinical examination, and case number of the horse. Histopathologic findings consistent with pathology were recorded and assigned as the final diagnosis for each horse.

**Statistics**

Statistical analyses were performed using R21 with the packages “lme4”22 for mixed models, FactoMineR for PCA23 and “ggplot2”24 for graphics. Reliability for each question was calculated according to Equations 1, 2, and 3 (Data S1) by fitting a mixed effects model with rating as the repeated measure outcome variable to determine the random effects of rater, horse, and time of assessment (live, video1, video2).25,26 The function “lmer” was used for dichotomous, ordinal, and continuous variables. Odds ratios (OR) were calculated and significance set using Fisher’s exact test and the package “epiR.”27 The intraclass correlation coefficient (ICC) was calculated as defined in formulas 1 to 4 (Data S1), all redefined.4,28,29

Agreement within assessors was measured as assessment-reassessment (test–retest, which is similar to intrarater assessment), with the assumption of no change in condition between ratings. The variation between assessments was primarily attributed to variation because of time of assessment. The variation between assessments was primarily attributed to variation because of time of assessment. The reliability for the modified ataxia-grading scale was good (ICC = 0.74) after the full neurologic gait assessment (Tables 2, 3). When the horses were split into 2 groups of either low (≤1/5, n = 12) or high grade (≥2/5, n = 13), the ICC for live scorings was 0.08 for the low-grade group and 0.43 for the higher grade group. When comparing agreement for the medicine group and the surgery group (Tables S2, S3), there is good and higher agreement for the surgery group on overall lameness scoring (ICC for medicine = 0.30 and ICC for surgery = 0.55) and a similar agreement on the overall ataxia scoring (ICC for both groups = 0.72). Clinicians often disagreed on grade of ataxia with the greatest disagreement over horses with a median grade of 2 (Fig 1).

The PCA revealed 2 principal components explaining 53% of the variation, with 41% of the first and 12% on the second dimension (Table 3). All gait-related assessments with an ICC ≥ 0.4 correlate significantly to the first principal component except “making mistakes over an obstacle,” “deficits within individual limbs” and “hopping,” all of which correlated with the second principal component (Table 4).

Sixteen horses were euthanized. Histopathologic changes consistent with pathology were detected in 7 ataxic horses: 1/3 horses with median ataxia grade 1, 4/6 horses with median ataxia grade of 2, and 2/3 horses with a median ataxia grade of 3. Histopathologic changes consistent with pathology identified in the ataxic horses ranged from classical Wallerian degeneration associated with compression (n = 6) to neuroaxonal dystrophy and a dorsal root neurofibroma (findings are summarized in Table 4). None of the 3 horses euthanized with a median ataxia grade of 0 had histopathologic changes consistent with pathology. Histopathologic evidence of disease in the brain, cerebellum or spinal cord was not identified in 5 horses with an ataxia grade ≥ 1; these included 1/3 with a median ataxia grade of 1, 2/6 with a median ataxia grade of 2, 1/3 with a median ataxia grade of 3, and 1/1 with a median ataxia grade of 4.

**Results**

The number of raters assessing each horse ranged from 3 to 6 with a median of 5 assessors. The group of horses comprised 12 mares and 13 geldings, with a mean age of 6.4 years, ranging from 3 to 16 years. Of the 25 horses, 7 had a median ataxia grade of 0; 5 had a median ataxia grade of 1; 6 had a median grade of 2; 6 had a median grade of 3; and 1 horse was graded as 4 (Table 1 and Table S4).

The reliability for the modified ataxia-grading scale was good (ICC = 0.74) after the full neurologic gait assessment (Tables 2, 3). When the horses were split into 2 groups of either low (≤1/5, n = 12) or high grade (≥2/5, n = 13), the ICC for live scorings was 0.08 for the low-grade group and 0.43 for the higher grade group. When comparing agreement for the medicine group and the surgery group (Tables S2, S3), there is good and higher agreement for the surgery group on overall lameness scoring (ICC for medicine = 0.30 and ICC for surgery = 0.55) and a similar agreement on the overall ataxia scoring (ICC for both groups = 0.72). Clinicians often disagreed on grade of ataxia with the greatest disagreement over horses with a median grade of 2 (Fig 1).

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**Components of the neurologic gait assessment considered abnormal that had an increased OR for spinal cord pathology included “walking with the head elevated,” “with a blindfold,” “hopping,” and “walking down a slope with the head elevated” (summarized in Table 4).**

Raters who knew that a horse was presented for evaluation of a neurologic or abnormal gait problem (excluding lameness) were more likely to assign a grade above the median ataxia grade (P = .02, mixed effect model) with an OR of 2.8 (95% CI = 0.8–9.5).

**Discussion**

Reproducible assessment of gait during a neurologic examination in horses is essential for the diagnostic
The most common neurologic diseases of horses affect the spinal cord, with resultant changes in gait caused by general proprioceptive deficits and paresis. In this study, the reliability for each part of the gait assessment ranged from poor to good; however, the only individual

### Table 2. Reliability results in the form of ICC for the dichotomous and categorical questions for gait assessment during the neurologic examination of horses. See questionnaire (Data S2) for full details on the questions and Tables S1, S2 for comparison of live to video and average rater agreement.

| Examination Parts | ICCa | Liveb | Video1 : Video2c |
|-------------------|------|-------|------------------|
| Walk and trot on a straight line | | | |
| Normal/abnormal | 0.40 | 0.43 |
| Neuro deficit type | 0.25 | 0.18 |
| Head elevation | | | |
| Normal/abnormal | 0.56 | 0.29 |
| Makes mistakes | 0.37 | 0.11 |
| Paretic | 0.35 | 0.31 |
| Ataxic | 0.55 | 0.17 |
| Hypermetric | 0.38 | 0.05 |
| Hypometric | 0.30 | 0.03 |
| Blindfold | | | |
| Normal/abnormal | 0.42 | 0.33 |
| Readily identifiable | 0.60 | 0.12 |
| Standing tail pull | | | |
| Normal/abnormal | 0.27 | 0.47 |
| Walking tail pull | Normal/abnormal | 0.37 | 0.25 |
| Small circles | | | |
| Normal/abnormal left | 0.48 | 0.34 |
| Normal/abnormal right | 0.44 | 0.39 |
| Circumducting | 0.45 | 0.30 |
| Turning normally FL | 0.12 | 0.00 |
| Turning normally HL | 0.21 | 0.12 |
| Backing up | Normal/abnormal | 0.69 | 0.47 |
| Limb placement | | | |
| Normal/abnormal LF | 0.46 | 0.66 |
| Normal/abnormal RF | 0.46 | 0.60 |
| Obstacle | Normal/abnormal | 0.47 | 0.48 |
| Makes mistakes | 0.68 | 0.67 |
| Mistakes are neurologic | 0.45 | 0.07 |
| Deficits LF | 0.56 | 0.09 |
| Deficits RF | 0.50 | 0.37 |
| Deficits LH | 0.52 | 0.70 |
| Deficits RH | 0.39 | 0.27 |
| Panniculus reflex | Normal/abnormal left | 0.37 | 0.67 |
| Normal/abnormal right | 0.24 | 0.61 |
| Cervicofacial reflex | Normal/abnormal left | 0.37 | 0.43 |
| Normal/abnormal right | 0.33 | 0.41 |
| Hopping | Normal/abnormal | 0.45 | 0.35 |
| Weight shift to pelvic limbs | 0.26 | 0.21 |
| Stumble | 0.55 | 0.34 |
| Asymmetric | 0.26 | 0.17 |
| Slope | Normal/abnormal | 0.42 | 0.19 |
| Mistakes | 0.32 | 0.09 |
| Paretic | 0.37 | 0.34 |
| Ataxic | 0.52 | 0.17 |
| Hypermetric | 0.26 | 0.15 |

### Table 2 (Continued)

| Examination Parts | ICCa | Liveb | Video1 : Video2c |
|-------------------|------|-------|------------------|
| Slope | Normal/abnormal | 0.42 | 0.19 |
| Mistakes | 0.32 | 0.09 |
| Paretic | 0.37 | 0.34 |
| Ataxic | 0.52 | 0.17 |
| Hypermetric | 0.26 | 0.15 |

### Table 3. Reliability results as ICC for the questions with answers on an ordinal scale. The ICCs are calculated from the gait assessment in the neurologic examination of horses. See questionnaire (Data S2) for full details on the questions and Tables S1, S2 for comparison of live to video and average rater agreement.

| Examination Parts | ICCa | Liveb | Video1 : Video2c |
|-------------------|------|-------|------------------|
| Walk and trot on a straight line | | | |
| Lame or neurologic | 0.21 | 0.09 |
| Lame leg | 0.49 | 0.35 |
| Ataxia grade | 0.71 | 0.48 |
| Lameness grade | 0.29 | 0.39 |
| Across all | Ataxia score LF | 0.39 | 0.28 |
| Ataxia score LH | 0.69 | 0.54 |
| Ataxia score RF | 0.37 | 0.28 |
| Ataxia score RH | 0.60 | 0.42 |
| Paresis score LF | 0.07 | 0.18 |
| Paresis score LH | 0.54 | 0.57 |
| Paresis score RF | 0.00 | 0.12 |
| Paresis score RH | 0.30 | 0.36 |
| Overall lameness grade | 0.26 | 0.39 |
| Overall ataxia grade | 0.74 | 0.59 |

ICC, intraclass correlation coefficient. Bolded numbers have an ICC ≥ 0.4 (fair agreement).

a ICC for a single rater (Equation 1, ICC1[A,1]).
b ICC live scoring only, single rater (Equation 1).
c ICC test-retest (video only), from 1st video session to 2nd video session (Equation 3).

process and for decision making in prepurchase examinations, for considering treatment options, safety for handlers and riders and animal euthanasia. The most common neurologic diseases of horses affect the spinal cord, with resultant changes in gait caused by general proprioceptive deficits and paresis. In this study, the reliability for each part of the gait assessment ranged from poor to good; however, the only individual
elements with an ICC > 0.6 were “backing up,” recognition of a horse “making mistakes over an obstacle,” overall ataxia score and pelvic limb ataxia score. The fair to poor agreement between raters and for assessment-reassessment has important implications for the daily clinical assessment of individual horses as well as follow-up examinations. In addition, there are implications for the assessment of neurologic gait deficits in research studies evaluating equine neurologic diseases and their treatments.

A perfect rating scale should be reliable, valid, responsive and enable identification of the clinically minimal important difference (CMID) between 2 treatment groups or in a change over time. Indeed, for decision making for individual patients, some authors recommend that the ICC should be at least 0.90. Based on these recommendations, neither the individual criteria for gait assessment within the equine neurologic examination, nor the modified ataxia-grading scale itself is acceptable for clinical use.

We found a higher overall agreement for live scoring of ataxia compared to previous studies assessing lameness scoring in a live setting using a 6-point scale (AAEP). In normal horses or those with subtle ataxia, agreement on the ataxia grading was worse than found in horses with low-grade lameness. Similarly, there was worse agreement between raters for their assessment of the moderate to high-grade ataxia group, compared to assessment of horses with higher grades of lameness. The relatively lower agreement for the moderate to high-grade ataxia could be explained by the large variation between raters when assessing horses with a median ataxia grade of 2 (Fig 1). We also found a more pronounced disagreement for the assessment of ataxia in horses compared to a new scale applied to assess dogs with spinal cord injury using the Texas Spinal Cord Injury Score (TSCIS) with separate components of gait, proprioceptive positioning and nociception. Two blinded raters had excellent agreement (ranging from 0.72 to 1.00) across all criteria when rating 36 dogs. However, the dogs used in the TSCIS study all had spinal cord dysfunction ranging from mild to severe and the study did not include any unaffected controls. Conceivably, some of the variation between raters in this study might have resulted from the confounding factor of concurrent musculoskeletal disease, as several horses in this study were considered to be lame. Although previous studies of agreement on lameness did not include assessment of ataxia, ideally all horses in this study would have also received a complete lameness investigation and diagnostic analgesia. Unfortunately, this was beyond the scope of our work, but it highlights the
importance of establishing better standards for gait assessment in horses with gait deficits caused by neurologic or musculoskeletal disease (or both).

In human medicine, clinicians developed the scale for the assessment and rating of ataxia (SARA) based on the neurologic examination. SARA has 1 underlying construct that explains 80% of the variation with 8 separate criteria, all scored on ordinal scales (gait, stance, sitting, speech disturbance, finger chase, nose-finger test, fast alternating hand movements, and heel-shin slide). SARA has high inter-rater reliability ICC (0.98) and a test–retest reliability ICC of 0.90. The equine modified ataxia scale in this study performed comparably to subscales of the ICARS, but poorly in comparison with the overall excellent agreement and reliability of SARA. However, only neurologists or senior neurology residents assessed the patients in the SARA work and it was conducted on greater numbers of subjects, meaning that a direct comparison with this study might be misleading.

A video-based study using an 11-point lameness scale revealed fair reliability (κ = 0.41) for agreement between 3 raters scoring lameness based on video (ranging from 0.30 to 0.58 for each pair of raters). Previous studies into agreement on assessment of lameness on the 6-point scale40–41 have a poorer agreement because they are video-based. In a study of objective kinematic assessment of ataxic horses, poor agreement between live and video scoring was attributed to the different conditions for live compared to video assessment; however, our results suggest that it was more likely caused by poor live agreement and low test–retest reliability.

For testing hypotheses, clinical signs should be measurable as either dichotomous (absent or present), categorical (such as absent, mild, moderate or severe) or as ordinal (scaled) variables. Scales with multiple, ordinal divisions are more sensitive to change than dichotomous scales32 and their use has higher reliability because of reduced random error. Furthermore, a scale’s validity is better assessed and improved when scores are assigned within multi-item scales.32 We therefore recommend that a group of experts gather to discuss and refine the observations of horses with neurologic gait deficits30 based on a series of standardized videos of horses with confirmed spinal cord disease. A multi-item scale with ordinal ratings and with simplified and reliable descriptions of clinical observations of gait could be the aim of such an expert panel and examination components in this study with relatively higher reproducibility could be used as a foundation. The aim would be a scale with improved sensitivity to change and sufficient detection of CMID in order to help discriminate the severity of neurologic gaits for determination of response to treatments and for informed decisions regarding prognosis and horse and rider safety.

We confirm a likely expectation bias in the gait assessment component of the neurologic examination, since raters who were aware that a horse was presented for an abnormal gait (excluding lameness) or a possible neurologic gait deficit, were more likely to assign an ataxia grade higher than the median grade. This finding is comparable to the expectation bias reported in lameness evaluation.7 Another source of

### Table 4. OR and PCA results for questions with an ICC > 0.4. OR results for a positive test with evidence of spinal cord pathology. The PCA shows correlation with each question of the 2 dimensions (D1 and D2).

| Examination Parts | PCA | OR + Pathology^a |
|-------------------|-----|------------------|
|                   |     | Corr D1^b | Corr D2^b |
| Walk and trot on a straight line |     |           |           |
| Normal/abnormal   | 0.0 | 0.00–52  | 0.82^e  | 0.07  |
| Head elevation    |     |           |           |
| Normal/abnormal   | ∞   | 0.65–∞   | 0.91^f  | −0.17 |
| Ataxic            | 7.0 | 0.33–417 | 0.67^g  | 0.07  |
| Blindfold         |     |           |           |
| Normal/abnormal   | ∞   | 0.26–∞   | 0.78^h  | −0.36 |
| Readily identifiable^c | 0.8 | 0.05–12  | 0.20    | −0.08 |
| Small circles     |     |           |           |
| Normal/abnormal   | 1.5 | 0.08–28  | 0.83^i  | 0.18  |
| Normal/abnormal   | 7.0 | 0.33–417 | 0.94^j  | 0.08  |
| Circumduction     | 5.0 | 0.28–294 | 0.73^k  | 0.26  |
| Backing up        | 3.0 | 0.15–188 | 0.70^l  | 0.05  |
| Limb placement    | 2.0 | 0.15–33  | 0.26    | −0.17 |
| Normal/abnormal   | 35.0| 1.2–1844 | 0.53^m  | −0.60 |
| Obstacle          |     |           |           |
| Normal/abnormal   | 1.7 | 0.12–23  | 0.84^n  | 0.03  |
| Makes mistakes    | 1.5 | 0.08–29  | 0.42^o  | 0.61^p |
| Mistakes are neurologic^d | 9.0 | 0.35–546 | 0.79^q  | −0.18 |
| Deficits LF       | 0.1 | 0–3      | −0.20   | 0.52^r |
| Deficits RF       | 1.1 | 1.1–22   | −0.09   | 0.31  |
| Deficits LH       | 0.3 | 0.01–7   | 0.48^s  | 0.62^t |
| Deficits RH       | 0.9 | 0.05–15  | 0.22    | 0.33  |
| Hopping           |     |           |           |
| Normal/abnormal   | ∞   | 0.91–∞   | 0.41^u  | −0.49 |
| Stumble           | 9   | 0.35–546 | 0.38    | −0.57 |
| Slope             |     |           |           |
| Normal/abnormal   | 1.5 | 0.08–29  | 0.83^v  | −0.23 |
| Ataxic            | 1.5 | 0.08–29  | 0.84^w  | 0.36  |
| Slope with head elevation |     |           |           |
| Normal/abnormal   | ∞   | 0.26–∞   | 0.78^x  | −0.31 |
| Ataxic            | 1.5 | 0.08–29  | 0.65^y  | 0.17  |

CI, confidence interval; OR, odds ratio; PCA, principal component analysis; ∞, the OR is infinitely high. Bolded numbers represent questions with a correlation of 0.5 or higher with that dimension of the PCA.

^aOR calculation where disease is considered histopathologic evidence of spinal pathology and exposure is a positive test during live assessment.

^bCorrelation of the question with the PCA derived first dimension (D1, 2) and second dimension (D2, 3).

^cIf abnormal, is the deficit readily identifiable? Yes or No answer.

^dIf making mistakes, the mistakes are likely to be a neurologic deficit?

^eSignificant correlation with that dimension.

^fSignificant OR on Fisher’s exact test.
bias is experience and training, \(^4^2\) where experts in lameness assessment are more consistent in their scoring than residents, interns, and students.\(^7,^4^3\) Excluding residents from our analysis did not improve the ICCs of the live assessment. In addition, the difference between reliability for medicine and surgery was minimal, although surgeons had a higher reliability for assessing lameness compared to the medicine group (Tables S3, S4).

In this study, the median score of all raters was considered as a horse’s true grade, and the variation across all raters was examined around this score (Fig 1). This system assumes that all raters have similar ability in identification of neurologic gait deficits, but this might not be true as experience likely varies considerably. Optimally, the raters’ scores would have been compared to a reference (“gold”) standard for disease severity and presence or absence of disease. We attempted this by examining rater ICCs for horses with and without histopathologic changes consistent with pathology. Spinal cord pathology was commonly associated with a perception of horses’ being abnormal either when “walking with the head elevated,” “blindfolded,” “hopping,” or when “walking on a slope”, though it was not possible to quantify or compare disease severity by histopathologic examination of the spinal cord. However, the low number of horses in this study reduces the power of these conclusions and the lack of pathologic changes in 5 of 12 euthanized horses with a median ataxia grade ≥1 limited our ability to evaluate fully the neurologic examination’s validity. Nonetheless, our results emphasize the not uncommon disparity between clinician and pathologist when assessing horses with neurologic gait deficits. Furthermore, histopathologic assessment suffers from similar caveats as clinical assessment with variation or error introduced by experience level of the pathologist or tissue artifacts. For example, incidental background findings (spheroids) are found at all stages and at all levels of the neuroaxis in horses without neurologic disease.\(^4^4\) In addition, assessment of the importance of neuropathologic changes can best be made when the pathologist is aware of the history and clinical signs,\(^4^5\) presumably by increasing pretest probability and to maximize the chances of sampling the affected areas. As such, the pathologist’s opinion of the significance of histopathologic changes in the context of a history or clinical signs is akin to a clinician’s use of history and other clinical information in assessing the significance of perceived deficits detected during subjective (neurologic) examination (ie, expectation bias being a positive discriminatory factor). Furthermore, the assumption that all horses with apparent neurologic gait deficits should have identifiable pathologic changes might well be flawed as dynamic functional deficits might occur with only intermittent spinal compression, as is believed to occur in humans.\(^4^6\) In addition, recent identification of an important genetic component in controlling gait in horses\(^4^7\) reveals the extent to which gait alterations can have a functional rather than pathologic basis.

We conclude that clinicians should be aware of poor agreement between skilled and experienced observers of gait abnormalities in horses and poor correlation between pathology and clinical signs. The agreement is worse when signs are mild, and clinicians should be cautious when making decisions about horses on the basis of a subjective assessment of gait during the neurologic examination, especially when signs are subtle. This is particularly important during prepurchase examination and when a decision might lead to euthanasia or retirement or is being made for insurance purposes. Similarly, clinicians should be cautious when drawing conclusions from an apparent change in a horse’s degree of neurologic compromise after management changes or treatments, particularly given the anamnesis bias we report. We suggest that the neurologic assessment of horses’ gaits could be improved by identification of a set of objective parameters that can quantify severity of ataxia in horses, ideally in a clinical setting.

Footnote

* HDC TM-700; Panasonic, Osaka, Japan

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

- **Data S1.** Materials and Methods.
- **Data S2.** Rater Questionnaire.
- **Table S1.** Reliability results in the form of intraclass correlation coefficient for the dichotomous and categorical questions for gait assessment during the equine neurological examination assessed by four to six expert raters. See questionnaire (Data S2) for full details on the questions.
- **Table S2.** Reliability results as intraclass correlation coefficient (ICC) for the questions with answers on an ordinal scale. The ICCs are calculated from the gait assessment in the equine neurological examination assessed by four to six expert raters during live sessions only. See questionnaire (Data S2) for full details of the questions.
- **Table S3.** Overview of the horses and their median ataxia grade, median lameness grade and conclusions from the spinal cord histopathology.
- **Table S4.** Level of clinical significance of the intraclass correlation coefficient ranges for agreement, after Cicchetti.³¹
- **Table S5.** Reliability results as intraclass correlation coefficient for the dichotomous and categorical questions in gait assessment in the equine neurological examination assessed by four to six expert raters during live sessions. See (Data S2) for full details of the questions.
- **Table S6.** Reliability results as intraclass correlation coefficient (ICC) for the questions with answers on an ordinal scale. The ICCs are calculated from the assessment of gait in the equine neurological examination assessed by four to six expert raters. See questionnaire (Data S2) for full details of the questions.