Multivariate Analysis and Machine Learning in Cerebral Palsy Research

Jing Zhang*

Department of Neurology, Washington University in St. Louis, St. Louis, MO, United States

Cerebral palsy (CP), a common pediatric movement disorder, causes the most severe physical disability in children. Early diagnosis in high-risk infants is critical for early intervention and possible early recovery. In recent years, multivariate analytic and machine learning (ML) approaches have been increasingly used in CP research. This paper aims to identify such multivariate studies and provide an overview of this relatively young field. Studies reviewed in this paper have demonstrated that multivariate analytic methods are useful in identification of risk factors, detection of CP, movement assessment for CP prediction, and outcome assessment, and ML approaches have made it possible to automatically identify movement impairments in high-risk infants. In addition, outcome predictors for surgical treatments have been identified by multivariate outcome studies. To make the multivariate and ML approaches useful in clinical settings, further research with large samples is needed to verify and improve these multivariate methods in risk factor identification, CP detection, movement assessment, and outcome evaluation or prediction. As multivariate analysis, ML and data processing technologies advance in the era of Big Data of this century, it is expected that multivariate analysis and ML will play a bigger role in improving the diagnosis and treatment of CP to reduce mortality and morbidity rates, and enhance patient care for children with CP.

Keywords: multivariate analysis, machine learning, cerebral palsy, early diagnosis, outcome assessment

INTRODUCTION

Cerebral palsy (CP) is the most common movement disorder in children (1) and causes the most severe physical disability in neurodevelopmental disorders (2). Among children with CP, around 33% of them can not walk, 25% can not talk, 25% have epilepsy, 50% have an intellectual disability, and most of them are in pain (3). Spastic CP is the most common subtype of this disorder, often shown as muscle stiffness that causes movement difficulties in a hand, arm, foot, or leg on one or both sides of the body, affecting the majority (>85%) of children with CP (4). Other subtypes of CP include dyskinetic (athetoid or dystonic) CP, ataxic CP (e.g., with tremors), and mixed CP.

Currently, there is no cure for CP, but medications (such as baclofen and botulinum toxin), supportive treatments (such as physical therapy), and surgical procedures [such as orthopedic surgery and selective dorsal rhizotomy (SDR)] can help patients alleviate symptoms and improve motor skills (5). The signs and symptoms of CP usually appear in the early months of life, but the average age for diagnosis of CP is around 2 years (2). Therefore, early identification and intervention is crucial for patients with CP because infants have higher potential for recovery from neural lesions than adults (1, 2). Neuroimaging, motor assessment (such as general movement assessment), and neurological examinations can help identify high-risk infants, monitor
neurodevelopment, and detect or predict CP. Neuroimaging such as magnetic resonance imaging (MRI) and cranial ultrasound are useful to detect structural changes [intraventricular hemorrhage, periventricular leukomalacia (PVL), etc.] in the newborn brain, monitor lesion progression, and assess treatment effects, although compared with MRI, cranial ultrasound is less sensitive to lesions in the gray matter or malformations. Severe CP (caused by severe brain lesions such as PVL) can be identified by MRI or cranial ultrasound as soon as the lesions become recognizable on imaging after birth. However, 12–14% of children with CP have negative MRI scans due to subtle lesions in the brain (6). Thus, an integrated approach (imaging, motor assessment and neurological examinations) is needed to predict mild or moderate CP.

To predict CP in infants, Prechtl has described a general movement assessment method as a clinical assessment approach to identify CP motor impairments in infants by evaluating their spontaneous general movements (7). In particular, two atypical motor development features [(1) the presence of cramped-synchronized general movements at a preterm or term age and (2) the absence of small smooth movements or fidgety movements at 3–5 months] have been defined (8–10), which can identify CP in high-risk infants reliably (11). However, only well-trained physicians can perform such assessment, and general movement assessment based on visual observation by physicians is often influenced by subjective impressions and observer fatigue. Therefore, there is growing interest in developing multivariate and machine learning (ML)-based movement assessment tools for a more objective and quantitative motor assessment to detect movement impairments in high-risk infants (12, 13).

Multivariate analysis is a statistical analytical approach that simultaneously evaluates multiple variables, which compared with univariate analysis, may have more advantages (e.g., free from restrictions of various assumptions in univariate analysis) in identifying the associations between multiple data variables (e.g., variables associated with CP outcomes), grouping data into different groups or subgroups (e.g., different CP subtypes), and developing new diagnostic tests (e.g., differentiate CP subtypes with key feature variables). Multivariate analysis includes statistical methods such as principal components analysis (PCA), canonical variate analysis, independent components analysis, and multivariate regression. ML (or statistical learning) is a group of multivariate analytic methods that first identify the most significant data features or patterns that can best separate the data into different classes in the training dataset, and then apply these data features or patterns to the test dataset for data classification or prediction. ML has been increasingly applied to the biomedical field (14, 15), and examples of ML methods include linear discriminate analysis (16), support vector machine (SVM) (17), artificial neural networks (ANN) (18), random forest (19), and cluster analysis (20).

With growing interests, multivariate analysis has been increasingly employed in CP research in recent years, and research with multivariate analyses in CP is in infancy (14). To provide an overview of this relatively young field, PubMed search was performed with keywords “multivariate analysis cerebral palsy pediatric,” “machine learning cerebral palsy,” or “multivariate analysis cerebral palsy imaging.” The search yielded 126 articles. Articles were excluded if their subjects were not pediatric or the statistical methods used were not multivariate or the article was published before year 1990. This paper assessed the studies that used multivariate analysis in CP research and found that multivariate studies in CP are mainly in four categories: (1) risk factor identification; (2) detection of CP and identification of CP abnormalities; (3) movement assessment for CP prediction; and (4) outcome evaluation.

**MULTIVARIATE ANALYSIS IN RISK FACTOR IDENTIFICATION**

Early work on CP risk factor identification started from birth certificates. In a large population-based cohort study, data from birth certificates for 192 children with CP in four counties in California were compared with 155,636 healthy children in the same regions and the study found that low birth weight and (early or late) gestational age at birth were associated with high prevalence of CP, but early prenatal care and delivery at a hospital (for low birth weight neonates) were not associated prevalence of CP (21). Using multivariate analysis on clinical data of 113 CP infants (identified from 1,105 infants), Pinto-Martin et al. found that in low birth weight infants, cranial ultrasound imaging abnormalities such as parenchymal echodensities/lucencies (or ventricular enlargement) and germinal matrix/intraventricular hemorrhage were strong risk factors for disabling CP, but factors such as birth weight, gestational age, and Apgar score were not associated with it (22). In addition, a multicenter, large sample study (across eight European study centers, n = 585) revealed that there was a high rate of infection in mothers of CP children during their pregnancy and major CP abnormalities on structural MRI included white-matter damage due to immaturity (e.g., PVL) (42.5%), lesions in the basal ganglia (12.8%), cortical or subcortical lesions (9.4%), and malformation (9.1%) (23). A number of studies that identified CP risk factors have performed both univariate and multivariate analyses (24–26) where the risk factors identified by multivariate analyses were a subset of those identified by univariate analyses (24, 25), and the results of multivariate analyses were more rigorous and valid.

Studies that applied multivariate analysis to CP risk factor identification are summarized in Table 1 (22, 24–31). Multivariate logistic regression has often been used to identify risk factors for CP (22, 26, 29–31). Some of the risk factors identified by multivariate analytic studies include: premature birth, low birth weight, severe birth asphyxia, preterm rupture of membrane, abnormal cranial ultrasound, or structural MRI imaging findings (e.g., parenchymal echodensities/lucencies, or ventricular enlargement), intraventricular hemorrhage, PVL, neonatal sepsis, hypoxia-ischemic encephalopathy, hypoglycemia, neonatal jaundice, etc. (22–31). These risk factors are useful to understand the causes of CP, identify high-risk infants, and aid in the diagnosis of CP.

Further, the risk factors for CP revealed by the multivariate studies are useful to prevent CP. Several CP risk factors such...
### TABLE 1 | Summary of studies with multivariate analyses in identification of risk factors and detection of CP.

| Study        | Subject sample | Data                                                                 | Methods                                                                 | Main findings                                                                 | Other findings                                                                 |
|--------------|----------------|----------------------------------------------------------------------|-------------------------------------------------------------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------|
| Pinto-Martín et al. (22) | 113 children with CP | Clinical data (birth weight, gestational age, length of hospital stay, gender, race, plurality, presence of labor, Apgar score, motor function, cranial US findings, etc.) | Multivariate logistic regression to assess risk factors for CP | Risk factors for disabling CP: PVL/VE or ventricular enlargement on cranial US, germinal matrix/intraventricular hemorrhage, mechanical ventilation; risk factors for non-disabling CP: PVL/VE | Cranial US abnormalities are strong risk factors for disabling CP in low birth weight infants; non-risk factors for disabling CP: birth weight, gestational age, length of hospital stay, gender, race, plurality, presence of labor, Apgar score |
| Allan et al. (24) | 36 pts with CP (in 381 infants) | Clinical data (birth weight, bronchopulmonary dysplasia, abnormal cranial US findings, treatment, etc.) | Univariate and multivariate analysis to identify antecedents of CP | Predictors of CP: bronchopulmonary dysplasia and an abnormal cranial US scan (showing grade 3 to 4 intraventricular hemorrhage, PVL, or ventriculomegaly) | PVL and ventriculomegaly associated with high CP detection rates; chorioamnionitis and treatment with surfactant significant in univariate analysis |
| Han et al. (27) | 21 children with CP | Clinical data (birth characteristics, disease at birth, neonatal cerebral ultrasound findings, etc.) | Multivariate analysis used to identify risk factors for CP | Risk factors for CP: existence of PVL, preterm labor, preterm rupture of membrane, severe birth asphyxia, neonatal sepsis, and respiratory distress syndrome | Existence of PVL is the strongest risk factor for CP |
| Zhong et al. (28) | 308 children with CP | Data from a cross-sectional survey (birth characteristics, disease during the first month of life, etc.) | Multivariate analysis used to identify risk factors for CP | Risk factors for CP: delivery at home, low Apgar score, illness during the first month of life, maternal cold with fever in early gestation, low protein intake during pregnancy, low education level of mother |  |
| Golomb et al. (26) | 76 children with CP after perinatal stroke | Clinical data (perinatal history, motor function, frequency of CP, degree of disability, etc.) | Univariate and multivariate analysis (with logistic regression) to assess risk factors for CP in perinatal stroke | 68% pts with perinatal stroke had CP; risk factors for CP: delayed stroke and male gender; In pts with neonatal stroke, risk factors for triplegia or quadriplegia: bilateral infants | In pts with unilateral middle cerebral artery infarcts, risk factors for CP: delayed stroke and large-branch infarction |
| Miamoto et al. (29) | 60 pts with CP vs. 60 healthy controls | Data from questionnaires and clinical exams (TMD symptoms, bio-psychosocial characteristics, etc.) | Multivariate logistic regression to determine risk factors for TMD symptoms | Risk factors for TMD symptoms: presence of CP, male gender, severity of the malocclusion, mouth breathing, and mixed dentition | 13.3% pts vs. 1.7% controls had TMD symptoms |
| Abdullahi et al. (30) | 111 pts with CP vs. 222 controls | Clinical data (maternal sociodemographic characteristics and neonatal expected predictors) | Univariate and multivariate (logistic regression) analyses used to identify factors associated with CP | Predictors of CP: maternal fever, previous neonatal death, and poor sucking | Factors not associated with CP: maternal age, parity, birth weight, and sex |
| Yu et al. (31) | 203 preterm infants with CP, vs. 220 preterm infants without CP or other neurological disorders | Data of diseases of premature infants, the treatments in neonatal period, etc. | Multivariate logistic analysis used to identify risk factors associated with CP | Risk factors for CP: occurrence of PVL, HIE, hypoglycemia, or neonatal jaundice | Continuous positive airway pressure may lower the risk of CP |
| Golomb et al. (32) | 76 children with CP after perinatal stroke | Clinical data (perinatal history, motor function, frequency of CP, degree of disability, etc.) | Univariate and multivariate analysis (with logistic regression) to assess association of CP with other disabilities | 72% pts with perinatal stroke had at least another disability; risk factors for epilepsy: neonatal presentation and history of cesarean-section delivery | Risk factors for severe cognitive impairments or epilepsy: perinatal stroke with neonatal presentation |

(Continued)
### TABLE 1 | Continued

| Study                | Subject sample | Data                                                                                           | Methods                                                                                      | Main findings                                                                                       | Other findings                                                                                           |
|----------------------|----------------|------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------|
| Griffiths et al. (33)| 20 pts with spastic CP; 20 with dyskinetic CP | Injury severity scores at different brain regions on magnetic resonance imaging (T2)        | Variables indicated by univariate analysis fed to multivariate logistic regression to identify predictors to differentiate spastic and dyskinetic CP | Spastic CP pts had more severe damage to white matter near the paracentral lobule; dyskinetic CP pts had more injury to the STN; hypoxic-ischemic injury to the STN at birth associated with dyskinetic CP | Non-predictors of dyskinesia: injuries to the putamen, caudate, and globus pallidus |
| Yoshida et al. (34)  | 34 pts with CP vs. 21 healthy subjects | Parameters (number of fibers, tract-based FA, and FA for CST and posterior thalamic radiation tracts from diffusion tensor imaging (DTI) and motor level data | Univariate and multivariate (regression) analysis used to identify variables correlated to gross motor function | Number of fibers and ROI-based FA values of both tracts were lower in pts than controls; motor-sensory parameters were negatively correlated with GMFCS level | In pts with CP, mental retardation and epilepsy, epilepsy is an aggravating factor on bone health |
| Coppola et al. (35)  | Group 1: 40 pts with CP and mental retardation; group 2: 47 pts with CP; mental retardation and epilepsy; group 3: 26 pts with epilepsy | Clinical data (age, BMI, BMD z-score from dual-energy X-ray absorptiometry scan, etc.) | Multivariate analysis used to identify factors on BMD | Lower BMD in 42.5% pts in group 1, 70.2% in group 2, 11.5% in group 3 | Factors on BMD: age, BMI, severe mental retardation, epilepsy |
| Benfer et al. (36)   | 120 pts with CP | Data of OPD measures, motor measures, etc.                                                      | Univariate and multivariate regression analysis to determine the relationship between OPD and motor functions | Higher odds of OPD in non-ambulant pts than in ambulant pts | 85% pts had OPD |
| Romeo et al. (37)    | 100 pts with CP (32 of them with epilepsy) vs. 100 healthy children | Data from the SDSC, GMFCS levels, etc.                                                          | Multivariate analysis (logistic regression) used to identify factors associated with SDSC        | 13% of children with CP had abnormal sleep score; factors associated with SDSC: behavioral problems and epilepsy | Compared with healthy controls, sleep disorders are more common in children with CP |
| Adler et al. (38)    | 18 children with unilateral spastic CP (9 with mirror movement, 9 without) | Clinical data from BANIMM, JTHFT, and AHA                                                       | Multivariate analysis of covariance used to determine whether mirror movements affect daily living | Mirror movements had a negative impact on bimanual performance (AHA) and on the time needed to complete difficult activities | Abnormal MMSE curve reflected problems in individual muscles such as motor control impairments, loss of muscle couplings, and spasticity or paralysis |
| Tao et al. (39)      | 11 children with CP, 8 healthy children, 7 healthy adults | EMG data from five thigh muscles and three lower leg muscles                                    | Multivariate empirical mode decomposition enhanced MMSE analysis used to analyze EMG data; repeated-measure ANOVA for group comparison | Compared with the control group, CP pts had distinct diversity in MMSE curve | Abnormal MMSE curve reflected problems in individual muscles such as motor control impairments, loss of muscle couplings, and spasticity or paralysis |
| Ghate et al. (40)    | 54 pts with CP | Clinical data (CP type, motor function, etc.) and data from ophthalmoscopic examinations      | Multivariate logistic regression to identify factors associated with motor outcomes             | 70% pts had abnormal optic nerve head disk; pialar associated with non-ambulatory status and quadiplegia; large cup associated with age at examination | Indicator for poor motor outcome: presence of optic nerve head pialar |
| Reid et al. (41)     | 31 children with unilateral CP | Activation maps from fMRI (with hand task); FA and MD values and fiber tracts in the thalamocortical and corticomotor tracts from DTI; clinical scores of motor ability | k-means clustering used to identify fMRI-task-specific DTI tracks; surface-based approach (using surface-meshes) compared with voxelwise fMRI-DTI approach; correlation analysis between DTI metrics and clinical scores performed | DTI metrics and five clinical scores of motor function were correlated; surface-based approach processed more subjects’ data (87%) than the voxel-based approach (65%), generated more coherent tractography | Surface-based approach revealed more significant correlations between DTI metrics and five clinical scores |
as brain injury and infection can be managed and avoided by preventing their causative mechanisms, and preventive efforts such as rubella vaccination, anti-D vaccination, and preventing methylmercury contamination are effective in preventing CP (2). In addition, meta-analysis has indicated that CP may be reduced by 30% in premature infants (<32 weeks gestation) by providing mothers of imminent labor with magnesium sulfate for neuroprotection of their babies (43, 44). Further, early interventions such as hypothermia have prevented CP in 12.5% of infants with neonatal encephalopathy following an acute intrapartum hypoxic event (45). Since, currently, there is no cure for CP, CP prevention is critical to reduce the prevalence of CP and save children from CP and CP-caused life-long disabilities. Multivariate analysis may help identify significant risk factors and early interventions in order to prevent CP.

Taken together, multivariate analysis is important in identification of risk factors for CP and the risk factors identified such as premature birth and abnormal (cranial ultrasound or structural MRI) imaging findings are useful not only for CP cause identification and diagnosis but also for CP prevention. Further research is needed to identify more manageable and avoidable risk factors and early interventions (such as neuroprotective drugs or therapies) to prevent CP and reduce CP morbidity rate.

**MULTIVARIATE ANALYSIS IN DETECTION OF CP AND IDENTIFICATION OF CP ABNORMALITIES**

Since the current average age for diagnosis of CP is around 2 years and infants have higher potential for neural recovery (1, 2), early detection of CP is critical to make early intervention possible. To identify CP in high-risk infants, neuroimaging such as cranial ultrasound and MRI is important for lesion detection and deciding the timing of the lesion. Studies applied multivariate analytic methods to detection of CP and identification of CP abnormalities are summarized in the latter part of Table 1 (32–42). A multicenter study of very-low birth weight infants (n = 381, survival rate = 87%, 36, or 9.4% with CP) indicated that cranial ultrasound findings such as grade 3–4 intraventricular hemorrhage and PVL were useful in predicting CP; in particular, PVL and ventriculomegaly were related to high detection rate (≥30%) for CP (24). Further, to differentiate CP subtypes, Griffiths et al. examined the T2 MRI images of children with spastic or dyskinetic CP (n = 20 in each group), and found that patients with spastic CP had more severe injury to white matter near the paracentral lobule, while patients with dyskinetic CP had more injury to the subthalamic nucleus (STN) (33). Multivariate logistic regression further identified the associated factors (i.e., lesions in distinctly different anatomical locations) for differentiation of spastic and dyskinetic CP (33).

When brain injuries in patients with CP are subtle, advanced imaging such as diffusion tensor imaging (DTI) and diffusion-weighted imaging is useful to detect CP injuries with subtle abnormalities. DTI metrics such as mean diffusion (MD) and fractional anisotropy (FA) are often used to identify injuries in white matter tracts. The value of DTI in identifying degenerative changes in patients with spastic CP due to periventricular white matter injury has been demonstrated by an early study, which reported that children with spastic hemiparetic CP (caused by periventricular white matter injury) had reduced DTI fiber count on the ipsilateral (the same side as the lesion) side of the corticospinal tract (CST), corticobulbar tract (CBT), and superior thalamic radiation, and had MD and FA changes reflected neurodegeneration of the motor and sensory pathways (n = 5) (46). Further DTI studies found that white matter damage in the posterior thalamic radiation pathways was more severe than that in the CSTs in children with CP (n = 28) (47), and DTI abnormalities in several white matter tracts such as posterior thalamic radiation pathways or superior regions of the thalamocortical and corticomotor tracts correlated with motor function measured by, e.g., Gross Motor Function Classification System (GMFCS) level (n = 28–34) (34, 41, 47). A review paper summarized the results of 22 DTI studies in CP and reported common findings of decrease FA (or increased MD) in the corticomotor and sensorimotor pathways, which correlated with clinical measures (48). Some research findings suggested that the CP injury in the somatosensory circuits might be more severe than that in the motor circuits, which may contribute more to motor impairment in CP (49). Further research is needed to unfold the mechanisms underlie sensorimotor impairment in CP and to improve detection of CP through neuroimaging.

Apart from neuroimaging, the abnormalities of CP have been identified via multivariate analysis using patients’ clinical data (perinatal history, CP type, CP frequency, motor function, etc.) to prevent CP and reduce CP morbidity rate.

**TABLE 1 | Continued**

| Study | Subject sample | Data | Methods | Main findings | Other findings |
|-------|----------------|------|---------|---------------|---------------|
| Tosun et al. (42) | 30 pts with CP only; 54 pts with CP and epilepsy, 38 pts with CP and high-risk children | BMD of lumbar vertebrae obtained by dual energy X-ray absorptiometry; clinical data (dietary Ca intake, whether intellectual disability, whether immobility, etc.) | Multivariate regression analysis used to evaluate the relationship between BMD and possible risk factors | Low BMD common in pts with CP and CP + epilepsy; risk factor of low BMD; immobility (not able to walk independently) | Low BMD related to the severity of CP, but not to vitamin D levels or AED treatment |

**AHA, assisting hand assessment; BANIMM, bimanual activities negatively influenced by mirror movements; BMD, bone mineral density; BMI, body mass index; CST, corticospinal tract; CP, cerebral palsy; EMG, electromyographic; FA, fractional anisotropy; GMFCS, Gross Motor Function Classification System; HIE, hypoxia-ischemic encephalopathy; JTHFT, Jebsen Taylor hand function test; MD, mean diffusivity; MMSE, multivariate multi-scale entropy; OPO, oropharyngeal dysphagia; PEJ/V, pentothal/sodium pentobarbital/lorazepam; Pts, patients; PVL, periventricular leukomalacia; ROI, region of interest; SDSC, Sleep Disturbance Scale for Children; STN, subthalamic nucleus; surgeon volume, the number of procedures performed; TMD, temporomandibular disorders; US, ultrasound.**
degree of disability, etc.) and data from other sources such as electromyography (EMG) and bone mineral density (BMD) (32, 35–40, 42). For example, multivariate analyses have indicated that 85% of CP patients have oropharyngeal dysphagia (36), 13% have sleep disturbance (37), 70% have abnormal optic nerve head (40), and 50% have low BMD (42). Factors associated with these CP abnormalities have also been identified (35, 37, 39).

In addition, since neonatal encephalopathy can cause CP, detection of neonatal encephalopathy helps detect potential CP. Structural brain connectivity networks of infants with neonatal encephalopathy have been examined using diffusion tractography extracted from DTI images, and ML methods such as SVM have been applied to structural connectivity features to detect neonatal encephalopathy (50). Moreover, since epilepsy and seizure disorders are common in children with CP, electroencephalography (EEG) is used to detect co-occurring seizures in high-risk infants or children. ML approaches such as SVM and ANN have been applied to EEG features to identify ictal and interictal spikes and achieved high detection rate for seizures (51). Further, multivariate analysis has found that children with CP after perinatal or neonatal stroke are more likely to have severe disability, cognitive impairment or epilepsy than CP children after delayed stroke (32).

Taken together, multivariate analytic studies in CP detection have identified imaging markers such as intraventricular hemorrhage and PVL on cranial ultrasound (24), injury to white matter near the paracentral lobule or to the STN on T2 MRI images (33), and injury in the CSTs and the posterior thalamic radiation pathways on DTI images (34, 41, 46, 47), which are useful in detecting CP and differentiating CP subtypes. Multivariate analyses have also identified CP abnormalities and their associated factors from non-imaging data (32, 35–40, 42). Further research is needed to identify biomarkers at the early stages of the disease to improve the diagnosis of CP, reduce diagnosis delay, and allow early identification and intervention for CP.

**ML IN MOVEMENT ASSESSMENT FOR CP PREDICTION**

Studies reported applications of ML in movement assessment for CP prediction are summarized in Table 2 (52–61). Early identification of motor impairments in high-risk infants enables early detection of CP. The two atypical movement features (related to the cramped-synchronized general movements and the absence of fidgety movements) in general movement assessment are strong predictors for CP diagnosis (8–10). Based on these key motor impairment features, their movement characteristics and associated movement variables have been identified to detect movement impairments in high-risk infants (53–55, 57–59). ML approaches have made it possible to analyze recorded movement data and identify motor impairments automatically.

As a pioneer study, Meinecke et al. analyzed the 3D movement data of infants (n = 22, seven with CP) from video recordings, extracted an optimal combination of movement features with cluster analysis, and identified CP motor impairments with quadratic discriminant analysis (overall detection rate: 73%) (52). Further, the characteristics of fidgety movements and associated movement measurements have been identified to distinguish infants with movement impairments from those without (53, 54, 58). For example, extracting motion features related to fidgety movements (such as motion distance and relative frequency) from video recordings with an optical flow-based method, Stahl et al. examined the motion patterns of 82 infants (15 with CP), applied SVM classifier to detection of CP movement impairments, and achieved a good classification accuracy (93.7 ± 2.1% with features of relative frequency; sensitivity: 85.3 ± 2.8%; specificity: 95.5 ± 2.5%) (58).

Apart from video recordings, other movement recording systems such as accelerometers and electromagnetic movement tracking system have been employed for movement assessment and CP prediction. Heinze et al. examined the general movements of a group of newborns and infants (n = 23, 4 with CP) with accelerometers, selected optimal (combined) movement features with genetic algorithm, classified CP motor impairments with a decision tree-based classifier, and obtained overall detection rates of 88–92% (55). To distinguish the gait patterns between patients with CP (n = 4), patients with multiple sclerosis (n = 4), and healthy controls (n = 12), Alaqtash et al. extracted gait features from 3D ground reaction force data, compared the gait patterns of the three groups, and applied nearest-neighbor classifier and ANN to gait feature classification, which led to overall classification accuracies of 85% (ANN, with a combination of gait features) and 95% (after optimizing the gait features to an optimal set of six gait features) (56). Moreover, using electromagnetic movement tracking recordings, Karch et al. studied the general movements from 63 infants (10 with CP), extracted movement features such as joint centers, and computed stereotype scores with dynamic time warping, yielding a high CP classification accuracy with stereotype score of upper limb movement (sensitivity: 90%; specificity: 96%) (57). For a review on movement recognition techniques in general movement assessment for CP prediction in high-risk infants, see Ref. (12).

In addition, multivariate and ML approaches have been used in the assessment of physical therapy, and the effect of orthotic devices such as ankle foot orthosis on CP patients (55, 56). To evaluate the quality of exercises in CP physical therapy, Parmar and Morris (n = 5) applied four classifiers (SVM, neural networks, AdaBoosted decision tree, and dynamic time warping) to movement feature (joint and angle data in time or frequency domain) classification to identify correct or wrong exercise, and found that among the four classifiers, AdaBoosted decision tree performed the best with high classification accuracies (94.68% for joint data; 90.3% for angle data) (56).

However, these machine-learning-based movement assessment studies are at the early stage of research. For example, the subjects in the study of Alaqtash et al. (56) were healthy subjects (n = 5), the movement data of wrong exercises (with errors) were simulated data, thus, the results were preliminary. Further research is needed to apply machine-learning methods to real movement data of patients with CP. In addition, the sample sizes of patients with CP in these machine-learning-based
### TABLE 2 | Summary of studies with multivariate analytic and machine learning approaches in movement assessment and outcome evaluation in CP

| Study                     | Subject sample | Data                                                                 | Methods                                                                 | Main findings                                                                 | Other findings                                                                                         |
|---------------------------|---------------|----------------------------------------------------------------------|-------------------------------------------------------------------------|------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------|
| Meinecke et al. (52)      | 22 infants (7 at risk of CP, 15 healthy) | 53 parameters extracted from recorded 3D movement data              | Cluster analysis based on Euclidian distances and quadratic discriminant analysis used to find the best combined parameters and separate at risk infants from healthy ones | Overall detection rate (using an optimal combination of 8 parameters): 73% (sensitivity: 1.00; specificity: 0.70) |                                                                                                      |
| Berge et al. (53)         | 14 infants with CP (who had four types of fidgety movements)       | Motion features (1D, 2D, and Wigner-Ville time-frequency virtue/feature) extracted from video recordings of movements | Periodicity (fidgety movements characterized by periodic patterns); principal components analysis (PCA) for data reduction; Pattern recognition (compare movement patterns in video with known visual patterns of fidgety movements) | ENIGMA (a software tool) can assess general movements and detect fidgety movements in CP pts           |                                                                                                      |
| Adde et al. (54)          | 30 high-risk preterm and term infants (13 developed CP in 5 years vs. 17 non-CP) | Movement variables (e.g., quantity of motion, and centroid of motion to identify fidgety movements) extracted from video recordings | Mann-Whitney U test; Logistic regression to identify CP predictors; ROC analysis to assess CP classification accuracy | 1/13 of pts had fidgety movements; predictor of CP; combined variable (centroid of motion STD, quantity of motion mean, quantity of motion STD); prediction accuracy of the combined variable: sensitivity: 85%; specificity: 88% | Combined variable had the highest prediction accuracy; ambulatory and non-ambulatory function was predicted correctly in 90% pts with CP |
| Heinzle et al. (55)       | 4 infants with CP, vs. 19 healthy infants                          | 32 features (including velocity and acceleration) extracted from measurement of accelerometers | Optimal parameter combinations selected by genetic algorithm; a decision tree-based classifier used to differentiate between pts’ and controls’ data | Overall detection rate: 88–92% for all measurements                                                                 | The low-cost movement disorder detection system based on accelerometers is applicable to CP diagnosis in newborns |
| Alaqtash et al. (56)      | 4 pts with spastic diplegic CP, vs. 4 pts with multiple sclerosis, vs. 12 healthy controls | Gait features extracted from 3D ground reaction force data          | NNC and ANN used to classify gait features into three groups; leave-one-out resampling | Classification accuracy (weighted average): 85% (using a combination of gait features); 96% (using an optimal set of six features) |                                                                                                      |
| Karch et al. (57)         | 10 infants with spastic CP, vs. 53 non-CP infants                  | Stereotypy score of limb movements extracted from electromagnetic movement tracking recordings | A multi-segmental chain model used to calculate the joint centers and joint axes; dynamic time warping used to compute stereotype scores; ROC analysis used to assess CP classification accuracy | CP classification accuracy using stereotype score of upper limb movement; sensitivity: 90%; specificity: 96% | Using stereotype score of leg movement could not distinguish pts from controls |
| Stahl et al. (58)         | 82 infants (15 with CP, 67 healthy)                               | Motion features (such as motion distance and relative frequency) extracted from video movement recordings | Motion features selected to identify fidgety movements; SVM used to classify pts from controls; 10-fold cross-validation for classifier validation | Classification accuracy: with features of relative frequency: 93.7 ± 2.1%; sensitivity: 85.3 ± 2.8%; specificity: 95.5 ± 2.5% | Classification with other features (absolute motion distance and wavelet coefficient) had lower accuracy |
| Kanemaru et al. (59)      | 145 preterm infants (16 developed CP by 3 years of age, vs. 129 normal) | 6 movement indices (average velocity of limb movement, number of movement units, kurtosis of acceleration, jerk index, etc.) extracted from video recordings | Fisher’s exact test and Mann-Whitney U-test to distinguish pts from controls | CP pts had higher jerk index in the legs (p < 0.01), average velocity of the arms (p < 0.05), and number of movement units of the arms (p < 0.05) than controls | Jerkiness of spontaneous movements in preterm infants at term age is useful for predicting CP |
| Wäyä et al. (60)          | 51 children with diplegic CP vs. 34 healthy controls             | Spatiotemporal gait data (physical properties, walking speed, etc.) | Multiple regression normalization and standard dimensionless equations used for data normalization; multiple regression normalization to identify the effects of AFO on gait in pts | Multiple regression normalization revealed difference in more spatiotemporal parameters in pts who walked with and without an AFO; after multiple regression normalization, most spatiotemporal parameters in pts with AFO became closer to those of controls | Multiple regression normalization may be useful in evaluating CP gait and gait classification |

(Continued)
| Study | Subject sample | Data | Methods | Main findings | Other findings |
|-------|----------------|------|---------|---------------|----------------|
| Parmar and Morris (61) | 5 healthy subjects (who did exercises correctly, and also mimic the errors/mistakes in exercise made by CP pts) | Features (joint positions, angles) in the time domain (also transformed to the frequency domain) extracted from 78 training samples and 47 testing samples of physical exercises video recording | 4 classifiers (SVM, NN, AdaBoosted decision tree, and DTW) used to distinguish good and erroneous exercises (in five schemas such as Blast-Off exercise in CP physical therapy) | Classification accuracy: 94.68% for AdaBoosted tree on joint data (in time domain); 90.99% for SVM on joint data (in frequency domain); 90.65% for SVM on joint data (in time domain); 90.3% for AdaBoosted tree on angle data (in time domain) | Classification accuracy: 90.13% for single-layer NN on joint data (in time domain); 87.63% for SVM on angle data (in frequency domain); 74.03% for DTW on angle data (in time domain) |
| Hemming et al. (62) | 4,007 children with CP | Data from five CP registers (birth characteristics, severity of prematurity, motor function, history of seizures, etc.) | Kaplan-Meier survival estimates performed; Multivariate proportional hazards model fitted for survival analysis | Death rate: -8%; rate of children who survived to 20 years of age: 85–94%; predictors of CP survival: The number and severity of impairment | Birth weight and socioeconomic status might have impact on survival in certain register regions |
| Kim et al. (63) | 174 children with spastic CP who underwent SDR | Clinical data (age at surgery, types of CP, history of prematurity, motor function, history of seizures, etc.) | Univariate and multivariate logistic regression used to identify factors associated with surgical outcome | 6.3% pts had a poor outcome; predictor of outcome: type of CP (diplegia, quadriplegia) | Preoperative diagnosis was a strong predictor; intellectual delay was significant only in univariate analysis |
| Golan et al. (64) | 98 pts with spastic CP who underwent SDR | Data from hospital charts and radiographic spinal studies (preoperative and postoperative) | Univariate and multivariate regression analyses used to identify risk factors for spinal deformity | Risk factors for spinal deformity: CP severity; ambulatory function; age at surgery; gender | Factors associated with a lower rate of hyperlordosis; younger age at surgery and male gender |
| Majnemer et al. (65) | 95 children with CP | Data from Child Health Questionnaire and Pediatric QOL Inventory (by pts and parents), and measurements (Impairments, activity limitations, etc.) | Multivariate analysis used to identify determinants of QOL | Indicators of physical well-being: motor and other activity limitations; predictors of social-emotional adaptation: family functioning, behavioral difficulties, and motivation | 47% pts had mild motor impairment |
| White-Koning et al. (66) | 500 children with CP (in 7 countries in Europe) | Data from the Kidscreen questionnaire (by pts and parents) | Multivariate analysis used to identify factors associated the differences in parents' and pts' reports | Factors associated with the differences in parents' and pts' reports: high levels of stress in parenting (negative influence), self-reported severe child pain | Pts' self-reports higher than parents' in 8 domains, lower in the finances domain, and similar in the emotions domain |
| Long et al. (67) | 71 pts with CP vs. 77 non-CP; all subjects underwent orthopedic surgery | Demographic, surgical, and medical data (intraoperative opioid dosing, postoperative ICU admission, postoperative oxygen desaturation, etc.) | Multivariate regression analysis used to determine intraoperative opioid dosing associated outcomes and other variables | CP pts received less intraoperative opioid than non-CP pts; predictors of postoperative ICU admission and postoperative oxygen desaturation: intraoperative opioid dosing | CP associated with decreased opioid dosing |
| Smits et al. (68) | 116 pts with CP | 3-year longitudinal data (motor function, intellectual capacity, etc.) | Univariate and multivariate analyses to investigate associations between the course of capabilities (e.g., in mobility) and CP-, child-, and family characteristics | Predictors of self-care: a model including level of gross motor function and intellectual capacity; predictors of mobility: a model only including level of gross motor function; predictors of social function: a model including level of bimanual function and paternal educational level | Greater increase in capabilities for higher level of functioning, except for level of paternal education |
| Sponseller et al. (69) | 204 pts with CP who underwent spinal fusion surgery (at 7 institutions) | Clinical data of patient, laboratory, and surgical characteristics | Univariate and multivariate regression analysis to identify factors associated with infection development | 6.4% patients developed deep wound infection; factors associated with deep wound infection: presence of a gastrostomy/gastrojejunostomy tube | Determinants of apparent clearance: body weight, a possible genetic factor, and age |
| He et al. (70) | 61 pts with spastic CP | Serial R- and S-baclofen plasma concentrations | Mixed-effects population model and a 2-compartment model used for population pharmacokinetics analysis of oral baclofen; a final multivariable model used to describe oral baclofen profiles | Mean population estimate of apparent clearance: F: 0.273 L/h/kg with 33.4% IV; apparent volume of distribution (Vss/F): 1.16 L/kg with 43.9% IV; average baclofen terminal half-life: 4.5 h | Determinants of apparent clearance: body weight, a possible genetic factor, and age |

(Continued)
movement assessment studies are small (n = 4–15), studies with large samples are needed to further verify and improve these machine-learning methods. Further, the performance of these machine-learning studies or classification systems may be improved by optimizing the data processing chains (feature extraction, feature selection, classification, and verification). For details on optimizing data processing chains to improve classification performance, see Ref. (79).

Recently, Marschik et al. proposed an integrated system called a fingerprint model that monitors the movement and

| Study | Subject sample | Data | Methods | Main findings | Other findings |
|-------|----------------|------|---------|---------------|---------------|
| Kato et al. (71) | 31 pts with CP and cervical myelopathy; 30 with CSM, all pts underwent posterior decompression surgery | Measurements of pedicle and placement of pedicle screws from CT scans | Multivariate analysis used to evaluate factors associated with the breach of cervical pedicle screws | 23% CP pts and 7% CSM pts had pedicle sclerosis; pedicle sclerosis associated with a higher risk of breach | |
| Kruisjens-Terstra et al. (72) | 92 pts (2 years old) with CP | Longitudinal data (type of CP, GMFCS level, intellectual capacity, whether epilepsy, etc.) | Multivariate analysis used to identify determinants of development of self-care and mobility activities | Determinants of development of self-care activities: GMFCS and intellectual capacity; determinant of development of mobility activities: GMFCS | Self-care and mobility activity changes were less favorable in pts with severe CP |
| Shore et al. (73) | 320 children with CP who underwent VDRO for treatment of hip displacement | Clinical data (Age, sex, GMFCS, preoperative radiography, use of botulinum toxin, surgical performance, surgeon volume, etc.) | Univariate and multivariate (Cox regression) analyses used to determine effects of the data variables on surgical success; Kaplan-Meier survivalship curve generated | 92% success rate for GMFCS levels I and II vs. 76% success rate for GMFCS level V; predictor of surgical success: soft-tissue release at VDRO | 37% surgical failure; predictors of surgical revision: younger age at surgery, increased GMFCS level, and lower annual surgical hip volume |
| Mo et al. (74) | 206 children with CP who underwent surgical scoliosis correction | Clinical data (age, motor deficits, seizure history, verbal communication, mental retardation, Hydrocephalus severity, etc.) | Univariate and multivariate logistic regression used to identify factors causing poor IONM signals | Predictors of poor IONM signals: PVL, hydrocephalus, encephalomalacia; predictors of no signals: moderate or marked hydrocephalus, encephalomalacia | Predictors of no motor signal: focal PVL, moderate or marked hydrocephalus, encephalomalacia; predictors of no sensory signal: moderate hydrocephalus |
| Greco et al. (75) | 56 children with spastic CP | Clinical and neurophysiologic data (age, gross motor function, laterality of motor impairment, injury location and MEP) | Univariate and multivariate logistic regression analyses used to identify predictors of tDCS responses | Predictors of good responses to tDCS (and gait training): MEP (for 6-min walk test and gait speed), and subcortical injury (for gait kinematics and gross motor function) | The interaction of MEP and brain injury location predicted the responsiveness of tDCS |
| Minhas et al. (76) | 1,746 pts who underwent orthopedic procedure (345 pts underweight, 952 pts normal weight, 209 overweight, 240 obese) | Clinical data (whether seizure, whether asthma, whether use steroid, surgical procedure, etc.) | Multivariate logistic regressions performed to evaluate the effect of BMI on complications | Risk factors for total and medical complications in spine, hip, and lower extremity procedures: underweight class | Weight was not associated with complications in tendon procedures; overweight and obesity not associated with increased risk for complications |
| Galarraga et al. (77) | 115 children with CP who underwent (hip, ankle, foot, etc.) surgery | Preoperative data (36 physical examination variables and gait kinematics) and surgery data | PCA data dimension reduction; multi-regression analysis used to predict postoperative lower limb kinematics | Based on the kinematic angle, mean prediction errors on test vary from 4° (pelvic obliquity and hip abduction) to 10° (hip rotation and foot progression) | Mean prediction errors are smaller than the variability of gait parameters |
| Mann et al. (78) | 128 pts with CP | Physical activity, physical, psychosocial and total QOL reported by parents, walking performance measured by a StepWatch device | Multivariate regression used to examine the relationship of physical activity and walking performance to QOL | Physical activity positively associated with physical and total QOL; walking performance positively associated with physical QOL | Participation level positively associated with psychosocial QOL |

**TABLE 2 | Continued**
speech-language development of new-born babies and infants at risk and automatically detects neurodevelopmental disorders such as CP by multidimensional data analysis and machine-learning approaches (13). Although it is challenging (in technical details), the fingerprint model enables neurological assessment of at-risk infants in an objective and quantitative manner and facilitates early detection of CP and other neurodevelopmental disorders, which may be the future direction of pediatric clinical practice.

**MULTIVARIATE ANALYSIS IN CP OUTCOME EVALUATION**

Although there is no cure for CP, currently, treatment effects and outcomes in CP patients have been studied extensively. Multivariate approaches have been applied to outcome assessment (including survival analysis) in CP, and Table 2 (the latter part) provides a summary of these studies (62–78). The majority of the outcome studies employed a two-step approach: first, univariate analysis is used to identify variables that are associated with outcome; second, multivariate analysis is used to further examine the variables indicated by the univariate analysis and identify outcome predictors. A large sample of children with CP \( n = 4,007 \) in UK were studied, and multivariate survival analysis indicated that the death rate was \( \sim 8, 85–94\% \) of the children survived to age 20 years old, and the best predictors of CP survival were the number and severity of impairments (62). The multivariate outcome studies in CP fall into three categories: (1) outcome evaluation of medication and supportive treatments; (2) surgical outcome evaluation; and (3) quality of life (QOL) evaluation.

**Outcome Evaluation of Medication and Supportive Treatments**

The effect of commonly used medication oral baclofen on children with CP has been assessed with a multivariate model for the population pharmacokinetics analysis \( n = 61 \), and it has been found that baclofen dosage based on body weight was appropriate to treat patients \( \geq 2 \) years old, and determinants of apparent clearance in these children included body weight, a possible genetic factor, and age (70). Plasticity (shown as increased FA in the CSTs on DTI and improved motor function measures) induced by combined therapy (botulinum followed by physiotherapy) in children with spastic quadriplegia \( n = 8 \) has been reported (80), while a later DTI study indicated that the addition of botulinum to physiotherapy did not influence the outcome at 6 months in children with spastic diplegic CP \( n = 18 \) (81). DTI has also been used to evaluate the motor function outcomes of hemiplegic CP patients after rehabilitation treatment and DTI measurements such as the fiber number and FA of bilateral CSTs were correlated with functional level of hemiplegia scale (82). The quality of exercises in CP physical therapy has been evaluated with several classifiers, and AdaBoosted decision tree obtained good detection rate of exercise errors (61). In addition, multivariate outcome evaluation of therapies such as transcranial direct current stimulation (tDCS) has been performed (75).

Greco et al. investigated the functional outcome of tDCS in children with CP \( n = 56 \), and multivariate logistic regression analyses identified that the presence of motor evoked potential was a predictor for walk test and gait speed, subcortical injury was a predictor for gross motor function, and both of them were predictors of motor function gain arise from tDCS combined with gait training in these patients (75). However, there are few outcome studies of medications and supportive treatments in CP using multivariate analyses, and multivariate analysis may play a bigger role in such outcome evaluation to reveal the true therapeutic effects of these treatments and their outcomes in CP patients.

**Surgical Outcome Evaluation**

A number of multivariate studies have investigated the outcomes of surgical procedures in CP (63, 64, 67, 71, 74–77). For example, the outcomes of SDR surgery have been studied in children with CP and outcome-associated factors have been identified (63, 64). Kim et al. examined factors associated with poor outcome of SDR surgery in pediatric patients with CP \( n = 174 \) using multivariate logistic regression, and found that the poor-outcome rate was 6.3%, and the type of CP disability (diplegia, quadriplegia, etc.) was the predictor of poor outcome after SDR surgery (63). Further, Golan et al. evaluated the risk of spinal deformity in children with CP after SDR surgery \( n = 98 \) and multivariate regression analysis identified several risk factors for spinal deformity: CP severity, ambulatory function, age at surgery, and gender (64). In addition, Long et al. investigated intraoperative opioid dosing and associated outcomes with multivariate regression in children with CP \( n = 71 \) who underwent orthopedic surgery and reported that less intraoperative opioid was administered to CP children than non-CP children, and intraoperative opioid dosing was the outcome predictor of postoperative oxygen desaturation and ICU admission (67). Shore et al. examined the surgical outcome of children with CP \( n = 320 \) who underwent proximal femoral varus deotation osteotomy (VDRO) for treatment of hip displacement, and found that success rate was 92% for GMFCS levels I and II vs. 76% for GMFCS level V, and multivariate analysis indicated that soft-tissue release during the VDRO procedure was the predictor of surgical success (73). The surgical outcome predictors identified by these studies are useful for outcome prediction for individual CP patients.

Complications after CP surgical treatments have also been investigated. Using multivariate regression analysis, Sponseller et al. studied deep wound infection after spinal fusion surgery at seven institutions in pediatric patients with CP \( n = 204 \) and found that 6.4% of patients developed deep wound infection following surgery, and presence of a gastrostomy/gastrojejunostomy tube was the factor associated with infection (69). In addition, Kato et al. investigated cervical spine in patients with athetoid CP who underwent posterior decompression surgery \( n = 31 \) and multivariate analysis showed that pedicle sclerosis associated with a higher risk of breach of cervical pedicle screws (71). Further, Minhas et al. evaluated the effect of body mass index class on complications after orthopedic surgery in children with CP \( n = 1,746 \) and multivariate
logistic regression analysis revealed that underweight status was the risk factor for complications in osteotomies and spine surgery (76). The risk factors identified by these studies are helpful to avoid the surgical complications and improve surgical treatments in CP.

In children with CP who underwent surgery, intraoperative neuromonitoring (IONM) often fail (failure rate 61%) (74). Mo et al. studied IONM in children with CP who underwent surgical scoliosis correction (n = 206) and multivariate logistic regression analysis revealed that PVL, hydrocephalus, and encephalomalacia were the predictors of poor IONM signals, while moderate or marked hydrocephalus and encephalomalacia were the predictors of no signals (74). Further, outcome prediction of CP surgical procedures has been explored in a recent study. Galarraga et al. examined children with CP who underwent (hip, ankle, foot, etc.) surgery (n = 115), and multi-regression analysis revealed that preoperative and surgical data could predict postoperative kinematics, and mean prediction errors (varying from 4° to 10°) were smaller compared with the variability of gait parameters (77). These results are encouraging because they indicated that the postsurgical kinematics of patients with CP could be predicted (relatively accurately with small mean prediction errors) using presurgical and surgical data, which allows an estimate of postsurgical outcome ahead of time.

QOL Evaluation

Quality of life in physical ability, intellectual ability, self-care, and other aspects of life is an important outcome in CP. Multivariate analysis has been frequently used to assess QOL in patients with CP (65, 66, 68, 72, 78), and factors associated with physical QOL and self-care have been identified. For example, a multivariate analysis on QOL data of infants with CP (n = 92) identified GMFCS and intellectual capacity as the associated factors of self-care activity development, and GMFCS as the associated factors of mobility activities development (72). Further, a recent multivariate analysis showed that physical activity was positively associated with physical and total QOL in patients with CP (n = 128), and walking performance was positively associated with physical QOL (78). The factors identified by these studies may improve the QOL of patients with CP.

Taken together, since there is no cure for CP yet, and the death rate of CP is high (~8%), there is much to do to improve the outcomes of CP, and multivariate analytic approaches may play a bigger role in meeting such clinical demands. Surgical outcome predictors and risk factors for complications in CP surgical treatments have been identified by a number of multivariate outcome studies (63, 64, 67, 69, 71, 73, 76), which are useful not only for outcome evaluation and prediction but also for avoiding complications and improving surgical treatments in CP. However, there are few outcome studies for medications and supportive treatments (such as physical therapy) in CP using multivariate analysis. Thus, further research is needed to evaluate the outcomes of medications and supportive treatments, and multivariate analysis may play a bigger role in such outcome evaluation to reveal the true therapeutic effects of these treatments and their outcomes in CP patients, and help improve the outcomes of these treatments for patients with CP.

SUMMARY

Multivariate analysis has been applied to several areas in CP research such as identification of risk factors for CP, detection of CP and identification of CP abnormalities, movement assessment for CP prediction, and outcome assessment. The studies reviewed in this paper have demonstrated that multivariate analytic and ML approaches have made it possible to analyze movement recordings and identify CP movement impairments automatically. In addition, outcome predictors for surgical treatments have been identified by multivariate outcome studies. To make the multivariate analytic and ML approaches useful in clinical settings, further research with large samples is needed to verify and improve these methods in CP detection, movement assessment, and outcome evaluation/prediction. As multivariate analysis, ML and data processing technologies advance in the era of Big Data, it is expected that multivariate analysis and ML will play a bigger role in improving the diagnosis and treatment of CP to reduce mortality and morbidity rates, and enhance patient care for children with CP.

AUTHOR CONTRIBUTIONS

JZ reviewed the multivariate analytic studies in cerebral palsy and wrote up the manuscript.

ACKNOWLEDGMENTS

This work was inspired by the clinical team led by Dr. Manish Shah at the University of Texas Health Science Center (in Houston) and affiliated hospitals who are dedicated to patient care for children with cerebral palsy. Proofread was kindly provided by Drs. Gary and Carla Brandenburger.

REFERENCES

1. Herskind A, Greisen G, Nielsen JB. Early identification and intervention in cerebral palsy. Dev Med Child Neurol (2015) 57(1):29–36. doi:10.1111/dmcn.12331
2. McIntyre S, Morgan C, Walker K, Novak I. Cerebral palsy – don’t delay. Dev Disabil Res Rev (2011) 17:114–29. doi:10.1002/ddr.1106
3. Novak I, Hines M, Goldsmith S, Barclay R. Clinical prognostic messages from a systematic review about cerebral palsy. Pediatrics (2012) 130:e1285–312. doi:10.1542/peds.2012-0924
4. Reid S, Carlin J, Reddough D. Distribution of motor types in cerebral palsy: how do registry data compare? Dev Med Child Neurol (2011) 53:233–8. doi:10.1111/j.1469-8749.2010.03844.x
5. NIH (National Institute of Neurological Disorders and Stroke). Cerebral Palsy: Hope through Research. (2017). Available from: https://www.ninds.nih.gov/Disorders/Patient-Caregiver-Education/Hope-Through-Research/Cerebral-Palsy-Hope-Through-Research#3104_2
6. Krageloh-Mann I, Horber V. The role of magnetic resonance imaging in elucidating the pathogenesis of cerebral palsy: a systematic review. Dev Med Child Neurol (2007) 49:144–51. doi:10.1111/j.1469-8749.2007.00144.x
7. Prechtl HFR. Qualitative changes of spontaneous movements in fetus and preterm infant are a marker of neurological dysfunction. Early Hum Dev (1990) 23(3):151–8. doi:10.1016/0378-3782(90)90011-7
8. Einspieler C, Prechtl HFR. Prechtl's assessment of general movements: a diagnostic tool for the functional assessment of the young nervous system. Ment Retard Dev Disabil Res Rev (2005) 11:61–7. doi:10.1002/mrd.20051
9. Einspieler C, Marschik PB, Bos AF, Ferrari F, Cioni G, Prechtl HFR. Early markers for cerebral palsy: insights from the assessment of general movements. Future Neurology (2012) 7(6):709–17. doi:10.2217/fnl.12.60
10. Spittle AJ, Orton J. Cerebral palsy and developmental coordination disorder
11. Zhang Y, Wu J, Wu K, Ren R, Hou G, Peng D, et al. Evaluation of risk factors associated with cerebral palsy in children of Leshan Prefecture, Sichuan: a case control study on 308 cases. Zhonghua Fu Chan Ke Za Zhi (2002) 37(4):323–6.
12. Miamoto CB, Pereira LJ, Paiva SM, Pordeus IA, Ramos-Jorge ML, Marques LS. Prevalence and risk indicators of temporomandibular disorder signs and symptoms in a pediatric population with spastic cerebral palsy. J Clin Pediatr Dent (2011) 35(3):259–63. doi:10.17796/jcpd.35.3.738875v74l1m1p22
13. Marschik PB, Pokorny FB, Saha C, Garg BP, Azzouz F, Williams LS. Association of cerebral palsy with other disabilities in children with perinatal arterial ischemic stroke. Pediatr Neurol (2007) 37(4):245–9. doi:10.1016/j.pediatrneurol.2007.06.003
14. Griffiths PD, Radon MR, Crossman AR, Zurakowski D, Connolly D. Anatomic localization of dyskinesia in children with "profound" perinatal hypoxic-ischemic injury. AJNR Am J Neuroradiol (2010) 31(3):436–41. doi:10.3174/ajnr.A1854
15. Xu T, Rong L, Wang Q, You Y, Fu JX, Kang LM, et al. Influence of neonatal diseases and treatments on the development of cerebral palsy in preterm infant. Sichuan Da Xue Xue Bao Yi Xue Ban (2013) 4(2):270–3.
16. Golomb MR, Saad A, Garg BP, Azzouz F, Williams LS. Association of cerebral palsy with other disabilities in children with perinatal arterial ischemic stroke. Pediatr Neurol (2007) 37(4):245–9. doi:10.1016/j.pediatrneurol.2007.06.003
17. Grinevich PS, KadonMR, Crossman AR, Zurakowski D, Connolly D. Anatomic localization of dyskinesia in children with "profound" perinatal hypoxic-ischemic injury. AJNR Am J Neuroradiol (2010) 31(3):436–41. doi:10.3174/ajnr.A1854
18. Yoshida S, Hayakawa K, Yamamoto A, Okano S, Kanda T, Yamori Y, et al. Quantitative diffusion tensor tractography of the motor and sensory tract in children with cerebral palsy. Dev Med Child Neurol (2010) 52(10):935–40. doi:10.1111/j.1469-8789.2010.03669.x
19. Coppola G, Fortunato D, Mainolfi C, Porcaro F, Roccaro D, Signorelli G, et al. Bone mineral density in a population of children and adolescents with cerebral palsy and mental retardation with or without epilepsy. Epilepsia (2012) 53(12):2172–7. doi:10.1111/j.1528-1167.2012.03639.x
20. Benfer KA, Weir KA, Bell KL, Ware RS, Davies PS, Boyd RN. Oropharyngeal dysphagia and gross motor skills in children with cerebral palsy. Pediatrics (2013) 131(5):e532–42. doi:10.1542/peds.2012-3093
21. Romeo DM, Brogna C, Musto E, Baranello G, Pagliano E, Casalino T, et al. Sleep disturbances in preschool age children with cerebral palsy: a questionnaire study. Sleep Med (2014) 15(9):1089–93. doi:10.1016/j.sleep.2014.05.008
22. Adler C, Berweck S, Lidzba K, Becher T, Staadt M. Mirror movements in unilateral spastic cerebral palsy: specific negative impact on bimanual activities of daily living. Eur J Paediatr Neurol (2015) 19(5):504–9. doi:10.1016/j.ejpn.2015.03.007
23. Tao W, Zhang X, Chen X, Wu D, Zhou P. Multi-scale complexity analysis of muscle coactivation during gait in children with cerebral palsy. Front Hum Neurosci (2015) 9:367. doi:10.3389/fnhum.2015.00367
24. Ghate D, Vadanarayanan V, Kamour A, Corbett JI, Kedar S. Optic nerve morphology as marker for disease severity in cerebral palsy of perinatal origin. J Neurosci (2016) 36:258–31. doi:10.1523/JNEUROSCI.0629-16.2016
25. Reid LB, Cuningham R, Boyd RN, Rose SE. Surface-based fMRI-driven diffusion tractography in the presence of significant brain pathology: a study linking structure and function in cerebral palsy. PLoS One (2016) 11(8):e0159540. doi:10.1371/journal.pone.0159540
26. Tosun A, Erişen Karaca S, Ünvar T, Yurekli Y, Yenisey C, Omurulu IK. Bone mineral density and vitamin D status in children with epilepsy, cerebral palsy, and cerebral palsy with epilepsy. Childs Nerv Syst (2017) 33(1):153–8. doi:10.1007/s00381-016-3258-6
27. Kayem G, Mandelbrot L, Haddad B. Use of magnesium sulfate in obstetrics. Gynecol Obstet Fertil (2012) 40(10):605–13. doi:10.1016/j.gyobfe.2012.08.005
28. Crowther CA, Middleton PF, Voysey M, Askie L, Duley L, Pryce PG, et al. Assessment of the protective benefits for babies of antenatal magnesium sulphate: an individual participant data meta-analysis. PLoS Med (2017) 14(10):e1002398. doi:10.1371/journal.pmed.1002398
29. Jacobs SE, Tarnow-Mordi WO. Therapeutic hypothermia for newborn infants with hypoxic–ischaemic encephalopathy. J Paediatr Child Health (2010) 46:568–76. doi:10.1111/j.1442-2058.2010.01880.x
30. Thomas R, Eysen M, Peeters R, Molenaers G, Van Hecke P, De Cock P, et al. Quantitative diffusion tensor imaging in cerebral palsy due to periventricular white matter injury. Brain (2005) 128(Pt 11):2562–77. doi:10.1093/brain/awh600
31. Hoon AH Jr, Shashinko EE, Nagae LM, Lin DD, Keller J, Bastian A, et al. Sensory and motor deficits in children with cerebral palsy born preterm correlate with diffusion tensor imaging abnormalities in thalamocortical pathways. Dev Med Child Neurol (2009) 51(9):697–704. doi:10.1111/j.1460-7474.2009.03368.x
32. Schick SM, Boyd RN, Rose SE. New insights into the pathology of white matter tracts in cerebral palsy from diffusion magnetic resonance imaging: a systematic review. Dev Med Child Neurol (2012) 54(8):848–96. doi:10.1111/j.1469-8779.2012.04332.x
