Abstract

Background
Independence and frequency are two distinct dimensions of participation in daily life. The gap between independence and frequency may reflect the role of the environment on participation, but this distinction has not been fully explored.

Methods
A total of 18,119 parents or primary caregivers of children with disabilities aged 6.0–17.9 years were interviewed in a cross-sectional nationwide survey with the Functioning Scale of the Disability Evaluation System - Child version (FUNDES-Child). A section consisting of 20 items measured the children's daily participation in 4 environmental settings: home, neighborhood/community, school, and home/community. Higher independence and frequency restriction scores indicated greater limitation of participation in daily activities. Scores for independence, frequency, and independence-frequency gaps were examined across ages along with trend analysis. ANOVA was used to compare the gaps across settings and diagnoses for children with mild levels of severity of impairment.

Findings
A negative independence-frequency gap (restriction of frequency was greater than that of independence) was found for children with mild to severe levels of impairment. A positive gap (restriction of independence was greater than that of frequency) was found for children
with profound levels of severity. The gaps became wider with age in most settings of children with mild impairment and different diagnoses. Widest negative gaps were found for the neighborhood/community settings than for the other three settings for children with mild to severe impairment.

Conclusions

Children’s participation and independence-frequency gaps depend not only on the severity of their impairments or diagnoses, but also on their age, the setting and the support provided by their environment. In Taiwan, more frequency restrictions than ability restrictions were found for children with mild to moderate severity, especially in the neighborhood/community setting, and increased with age. Further identification of environmental opportunities that positively impact frequency of participation is needed.

Introduction

Developmental disabilities have been defined as a collection of chronic conditions in childhood and are manifested in cognitive, physical (e.g., motor, sensory), speech, language, or psychological impairments during the developmental period (birth up to 22 years) [1–3]. The launch of the International Classification of Functioning, Disability and Health: Children and Youth Version (ICF-CY) covers the important developmental period of childhood encompassing children’s disabilities and limited functioning of body functions/structures, activities, and participation. Its universal language and coding system can assist clinicians, educators, researchers, administrators, policy makers and parents to document and measure the important growth, health and developmental characteristics of children and youth [4].

Population-based nationwide surveys for children with developmental disabilities, usually enrolling children aged 11 to 22 years [3,5–9], play an important role in documenting epidemiological data, planning of services, evaluating prevention effectiveness and comparing health information. In Taiwan, the Functioning Scale of the Disability Evaluation System—Child version (FUNDES-Child) protocol was developed in conjunction with the launch of the People with Disabilities Rights Protection Act in 2007 [10]. The FUNDES-Child [11–13] is based on the Child and Family Follow-up Survey (CFFS) [14,15] for children and youth aged 6.0–17.9 years. The FUNDES-Child was developed to identify children with developmental disabilities in a nationwide pediatric population to prepare for the implementation of a new service policy based on the components of the ICF-CY.

The overall prevalence of disability or chronic conditions in children reported in nationwide surveys varies widely. For example, prevalence was estimated to be 1.22% in Taiwan [16], 17% in U.S. [7] and up to 30% in Canada [5]. Varying definitions of developmental disabilities is a major contributor to the variation of disability prevalence across countries. In the earlier definition in Taiwan, developmental disability was attributed mainly to health conditions or impairment of body functions, likely to be lifelong in nature and to result in substantial activity limitations and restrictions in societal participation [17]. In the new Disability Eligibility Determination System in Taiwan, rules [11,18] were adopted in which both impairment of body function and a diagnosis of disability are required to be eligible for services. In the US, diagnoses serve as the criteria for disability [7], whereas in Canada, the definition of developmental disabilities was broadened to cover chronic physical health conditions [5].
However, severity of body function impairments and diagnoses can only partly reflect participation restriction of children in daily activities [5,6,19]. The framework of the ICF-CY describes a dynamic interaction between the components of health conditions, body functions/structures, environmental and personal factors that contribute to children’s participation [4].

Based on the United Nations’ Convention on the Rights of the Child, participation has been described as the child’s right to be respected [20] and is considered an ultimate outcome for rehabilitation services based on the framework of ICF/ICF-CY [4,21]. Participation is known to be context-dependent as it describes the functioning of an individual’s social role. Therefore, settings, such as home, school, and community where children participate in daily activities should be identified in the investigation of children’s participation patterns. Recently, researchers have found that children with developmental disabilities face various environmental barriers that restrict their participation in home, school and community contexts [22–28].

In the ICF model (parent to the ICF-CY), Activity is defined as the execution of a task or action by an individual and Participation is defined by involvement in a life situation [4]. The component of Activity and Participation contains two constructs, “Performance” and “Capacity” [4]. However, participation of children has been defined with varied terms and multiple perspectives across measurement tools [29]. The construct of “Performance” describes what an individual “actually does or does do” in his or her current environment. In other words, “Performance” is viewed as “life experience” and is highly context dependent [4]. On the other hand, the construct of “Capacity” is defined as the full ability of an individual assessed in a standardized environment or uniform environment to neutralize the impact of different environments on the ability of the individual [4]. As such it may define what the child “can do in an ideal environment”, usually presented with the basic ability of developmental domains in children. Therefore, the “Capacity” construct identified in ICF is unsuitable to measure “Participation” because “Participation” refers to involvement in a “life situation” rather than in a “standardized” or “uniform” environment.

There is an array of measurable dimensions under the construct of participation in recently developed measures [29]. To explain the abilities in life situations, some researchers proposed the term “Capability” as a construct to clarify and operationalize the measurable dimensions of children’s “independence” or “range of possible functioning” in participation [29–31]. Morris defines capability as “the child’s predicament in the life they lead, taking into account their capacity and available resources, social and physical environment”[32]. Therefore, capability refers to what the child “can do in real life” in contrast to performance as “does do in real life”. In the FUNDES-Child, “capability” or “independence” describes the children’s abilities expected in life situation or in achieving a task or activities as assessed by a caregiver [11,12,14].

Another measurable dimension of participation used in the FUNDES-Child is “frequency”, referring to the extent to which a specific task or activity was carried out [29,33–35]. In this study, “Independence” refers to what a child “can do in real life” in contrast to “Frequency” which describes what a child “does do in real life”. The FUNDES-Child was designed to capture both independence and frequency of children’s participation in daily life as measured by the perspective of parents or primary caregivers [7–9,11–13,36,37].

The gap between “independence” and “frequency” is presumed to reflect the role of contextual facilitators and barriers on functioning and participation in that the environment often provides the context for the difference in what the child “can do” and “does do” [4,21]. However, there has been no clear evidence to support the associations between the independence-frequency gap and other ICF-CY components (health conditions, body functions, and contextual factors). Furthermore, children of school age face a transition from preschool to school settings, or from home to community contexts, a process that generates new life experiences. For children with disabilities, functional change across the life span, particularly in
participation, is critically important for adaptation and could be reasonably regarded as more important than the emergence of basic developmental skills. This distinction is important because the patterns of participation for varying age groups in population-based surveys should be taken into account when making social service policy to identify children’s needs and potential policy interventions.

Participation patterns in varying settings and across ages provide guidance for parents to describe their children’s needs in daily life. The concept of an independence-frequency gap can help in the development of policies and strategies to meet the needs of the children and their families. The purpose of this study was to explore the patterns of gaps in participation between independence and frequency as a function of severity of body functions and structures, diagnosed health conditions, settings and age.

Materials and Methods

This study utilized a national disability register based on a cross-sectional design of nationwide data collection in Taiwan. This national disability register provided the database of comprehensive profiles of participation in children aged 6.0–17.9 years. A 3-step data analysis was applied. Step 1 used all enrolled children covering a broad range of children to show the global profile of independence and frequency across age and setting in different severity groups. To control for possible confounders, Step 2 limited the children to the ones who had only one of the most frequent diagnoses to examine the effects of severity, setting and diagnosis on the independence-frequency gaps. Step 3 narrowed from Step 2 to include only children with mild level of impairment because of its large sample size that would provide adequate statistical power for each of the 12 age subgroups for trend analysis of frequency, independence and gaps in different diagnosis.

The study population

This study sample consisted of parents or caregivers of 18,119 children aged 6.0–17.9 years, who were interviewed between July 2012 and January 2014 following the official launch of the Disability Evaluation System (DES) [11,18,38]. The DES is a three-stage evaluation process: (1) medical examination (body functions/structures), (2) functional assessment (participation and environment), and (3) needs assessment [11,38]. Demographic data and health characteristics of the sample are presented in Table 1. The mean ages for the three data analysis steps were 12.1 (SD = 3.5) years, 12.0 (SD = 3.4) years, and 11.9 (SD = 3.4) years respectively. The numbers of children for each year age group were evenly distributed except for a higher percentage in the 12-year old group. The five largest diagnostic groups were intellectual disability (ID), autistic spectrum disorder (ASD), language delay, cerebral palsy (CP), and hearing impairment. The five diagnostic groups were collapsed into four categories in terms of functional limitations [1–3]: cognitive impairment (ID group), psychological impairment (ASD group), language/hearing impairments (LH group), and physical impairments (CP group) for statistical analysis at Steps 2 and 3 in this study.

Ethics approval

Ethics approval was obtained from Taipei Medical University- Joint Institutional Review Board. This study utilized the de-identified database of Taiwan Databank of Persons with Disabilities (TDPD). The children were assigned a diagnosis with specific codes of the International Classification of Disease, 9th Revision, Clinical Modification (ICD-9-CM) (http://www.cdc.gov/nchs/icd/icd9cm.htm) to be eligible for the DES.
Table 1. Demographic data for children for the three steps of data enrollment.

| Variables | Step 1 for all the enrolled children (N = 18,119) N (%) | Step 2 for children having only one type of the five diagnoses\(^a\) (N = 13,906) N (%) | Step 3 for children with mild severity having only one type of the five diagnoses\(^a\) (N = 7,719) N (%) |
|-----------|------------------------------------------------------|--------------------------------------------------|--------------------------------------------------|
| Sex, male n (%) | 11,698 (64.6) | 9,099 (65.4) | 5,187 (67.2) |
| Age band (Years) | | | |
| 6.0–6.9 | 1724 (9.5) | 1336 (9.6) | 719 (9.4) |
| 7.0–7.9 | 1334 (7.4) | 1009 (7.3) | 598 (7.7) |
| 8.0–8.9 | 1383 (7.6) | 1061 (7.6) | 615 (8.0) |
| 9.0–9.9 | 1397 (7.7) | 1067 (7.7) | 605 (7.8) |
| 10.0–10.9 | 1159 (6.4) | 888 (6.4) | 518 (6.7) |
| 11.0–11.9 | 1598 (8.8) | 1250 (9.0) | 754 (9.8) |
| 12.0–12.9 | 2036 (11.2) | 1622 (11.7) | 922 (11.9) |
| 13.0–13.9 | 1352 (7.5) | 1030 (7.4) | 554 (7.2) |
| 14.0–14.9 | 1544 (8.5) | 1209 (8.7) | 644 (8.3) |
| 15.0–15.9 | 1749 (9.7) | 1363 (9.8) | 668 (8.7) |
| 16.0–16.9 | 1377 (7.6) | 1022 (7.3) | 549 (7.1) |
| 17.0–17.9 | 1466 (8.1) | 1049 (7.5) | 573 (7.4) |
| Major diagnoses/ICD-9-CM codes\(^b\) | | | |
| Intellectual disability/317–319 | 10,310 (56.9) | 9,252 (66.5) | 4,966 (64.3) |
| Autism spectrum disorders/299.01–299.90 | 4,013 (22.1) | 3,331 (24.0) | 2,117 (27.4) |
| Language delay/315.31–315.39, 318.1 | 801 (4.4) | 497 (3.5) | 326 (4.2) |
| Cerebral palsy/343.9, | 602 (3.3) | 413 (3.0) | 105 (1.4) |
| Hearing impairment/389 | 448 (2.5) | 413 (3.0) | 205 (2.7) |
| Schizophrenia/295.10–295.90 | 191 (1.1) | 0 | 0 |
| Visual impairment/369.0–369.9 | 89 (0.5) | 0 | 0 |
| Cerebral vascular accident /431–438 | 58 (0.3) | 0 | 0 |
| Depression/296 | 31 (0.2) | 0 | 0 |
| Spinal cord injury/952,806 | 20 (0.1) | 0 | 0 |
| Impairment severity | | | |
| Mild | 9,561 (52.8) | 7,719 (55.5) | 7,719 (100.0) |
| Moderate | 5,614 (31.0) | 4,430 (31.9) | 0 |
| Severe | 1,774 (9.8) | 1,129 (8.1) | 0 |
| Profound | 1,170 (6.5) | 628 (4.5) | 0 |
| Occupation | | | |
| Student | 17,308 (95.5) | 13,520 (97.3) | 7,582 (98.1) |
| Employee | 33 (0.2) | 20 (0.1) | 12 (0.2) |
| Quit job/ drop out of school for health reasons | 485 (2.7) | 181 (1.3) | 45 (0.6) |
| Quit job/ drop out of school for non-health reasons | 141 (0.8) | 93 (0.7) | 44 (0.6) |
| Others | 152 (0.8) | 92 (0.6) | 36 (0.5) |
| Child’s physical health in general | | | |
| Excellent | 1,652 (9.1) | 1,407 (10.1) | 840 (10.9) |
| Very good | 3,894 (21.5) | 3,187 (22.9) | 1,836 (23.8) |
| Good | 4,756 (26.2) | 3,733 (26.8) | 2,113 (27.4) |

(Continued)
Measurement of body function

The severity of body functions/structures was assessed in the first stage of the DES. Relevant ICF/ICF-CY categories for specific diagnoses were coded by physicians trained in using a qualifier scale from 0 to 4 points presenting no problem (0), mild (1), moderate (2), severe (3), and profound (4). A final severity level of body function was assessed on the basis of decision rules for combining levels of severity of individual codes of body functions/structures. In this sample, half of the children were classified at mild levels of severity (Table 1).

Measurement of participation

Child participation was assessed using the FUNDES-Child Section II that contains 20 items to measure 2 dimensions, frequency and independence of participation [12]. The items from the Independence dimension were translated and modified from the Child and Adolescent Scale of Participation (CASP) [15], one part of the CFFS. The Frequency dimension has been designed by the Taiwan ICF team and added to each item of the FUNDES-Child Section II [11]. Items are scored by the 4 domains describing settings (home with 6 items, neighborhood and community with 4 items, school with 5 items, and home/community living activities [HCLA] with 5 items)[12].

The four domains present categories of activities children would experience in the four settings. Activities in the home setting include social, play or leisure activities, chores, self-care activities, communication and moving around at home; community setting activities include social, play or leisure activities, structured events, moving around and communicating with others in community; school setting activities include educational (academic) activities, social, play or leisure activities, moving around, and using educational material in schools; HCLA activities include household tasks, shopping and managing money, managing schedule, using transportation to get around, and work activities and responsibility in home and in transition to community. The following are examples of items for specific settings: “Communicating with other children and adults at home” (home), “Social, play or leisure activities with friends in the

Table 1. (Continued)

| Variables                          | Step 1 for all the enrolled children \(N = 18,119\) N (%) | Step 2 for children having only one type of the five diagnoses\(^a\) \(N = 13,906\) N (%) | Step 3 for children with mild severity having only one type of the five diagnoses\(^a\) \(N = 7,719\) N (%) |
|------------------------------------|----------------------------------------------------------|-----------------------------------------------------------------|----------------------------------------------------------|
| Fair                              | 6300 (34.8)                                              | 4665 (33.6)                                                     | 2521 (32.6)                                              |
| Poor                              | 1508 (8.3)                                               | 913 (6.6)                                                      | 408 (5.3)                                                |
| Missing                           | 9 (0.1)                                                  | 1 (0.0)                                                        | 1 (0.0)                                                  |

Child’s emotional health and well-being

|                  |               |                                               |                                                                    |
|------------------|---------------|-----------------------------------------------|--------------------------------------------------------------------|
| Excellent        | 738 (4.1)     | 555 (4.0)                                     | 335 (4.3)                                                          |
| Very good        | 2223 (12.3)   | 1658 (11.9)                                   | 913 (11.8)                                                        |
| Good             | 3995 (22.0)   | 3052 (21.9)                                   | 1733 (22.5)                                                       |
| Fair             | 7718 (42.5)   | 6044 (43.5)                                   | 3360 (43.6)                                                       |
| Poor             | 3434 (19.0)   | 2594 (18.7)                                   | 1376 (17.8)                                                       |
| Missing          | 11 (0.1)      | 3 (0.0)                                       | 2 (0.0)                                                           |

\(^a\) The five types of diagnoses are intellectual disability, autism spectrum disorders, language delay, cerebral palsy, and hearing impairment

\(^b\) Some children have more than one type of diagnosis at Step 1

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neighborhood and community” (neighborhood and community), “Using educational materials and equipment that are available to other children in his or her classroom that have been modified for your child” (school), and “Shopping and managing money” (HCLA) [40].

The FUNDES-Child utilizes a proxy format in which parents or caregivers answer questions about their child’s activities in the previous 6 months. In keeping with the format used in the FUNDES-Adult interview [11,38], flash cards with scoring options were used to assist parents in answering questions.

In the training manual and video of the FUNDES-Child, independence was defined as the child’s current level of ability to participate compared to that of other children of his or her age in the same community. For each item, independence was rated as: 0 (independent), 1 (with supervision/mild assistance), 2 (with moderate assistance), 3 (with full assistance). Frequency of participation was rated with reference to age as: 0 (the same or more than expected for age), 1 (somewhat less than expected for age), 2 (much less than expected for age), and 3 (never does). A response of not applicable (a child would not be expected to do that activity as peers of the same age and in the same community) was allowed for both dimensions. All items were rated under the condition that children used assistive devices as usual. As each item was on the same ordinal scale with the same anchor points at the extreme end (0–3 points), the two dimensions were comparable based on age-expected independence and frequency.

Items rated as “not applicable” were omitted in the scoring [40]. The mean scores for each of the 4 settings of FUNDES-Child Section II are thus the sum of scores all “applicable” items divided by the number of applicable items and then converted to a 0–100 scale. The certified testers could therefore interpret the scores within the same directional framework (higher scores represented greater restriction).

Training of FUNDES testers

Certified testers associated with 255 DES hospitals in Taiwan administered the FUNDES-Child by interviewing children’s parents or caregivers [41]. The certified testers were professionals licensed as physical therapists, occupational therapists, speech therapists, social workers, clinical psychologists, counseling psychologists, nurses, audiologists, special education teachers, and vocational evaluators. The training programs for certified interviewers covered the procedures of the DES and regulations (30 minutes), introduction to ICF and ICF-CY (30 mins), introduction to assessment instruments (FUNDES-Adult [60 mins], FUNDES-Child [40 mins]), practice of assessment instruments (200 mins), and the web-based platform for entry and storage of data (30 mins). At the end of each training course, a paper-and-pencil test was administered to certify the attending professionals [41].

Data collection

Children with disabilities were identified and recruited when they entered the DES system accompanied by his/her caregiver or parents. The first step was usually a visit with one physician followed by the FUNDES-Child assessment with a certified tester in a room in an authorized hospital. The interview and rating of the FUNDES-Child lasted about 40–60 minutes.

Data entry and analysis

To ensure data quality, all the assessment records were entered by the officers in hospitals into the TDPD, a nationwide web-based information system, and checked by the officer of the local department of health. The physician and the FUNDES qualified tester who completed the disability assessment report were required to enter their names into the information system. The
self-detection mechanism of the information system prevented data errors and missing responses. The data were then exported to statistical packages periodically.

According to the flow of the three steps of data reduction and analysis (Fig 1), the strategies of analysis are described below.

In Step 1, we included all the enrolled children (n = 18,119) to illustrate the global picture of the changes of independence, frequency and the gaps between them with age as a function as severity and settings. The mean scores for independence and frequency were plotted on dual Y coordinates (from age expected to most restricted based on a 0–100 scale), indicating the discrepancy between independence and frequency grouped by severity and settings across age (Fig 2).

In Step 2, we included the children seen only in one of the more frequent diagnostic groups (n = 13,906) to avoid the issue of comorbidity. The independence-frequency gaps were defined as the scores of independence restriction minus the scores of frequency restriction. If the independence-frequency gap was positive, it meant that restriction of independence was greater than that of frequency (or doing frequency exceeded doing ability); if the gap was negative, it meant that restriction of frequency was greater than that of independence (or doing ability exceeded doing frequency). The gap, discrepancy between independence and frequency, is wider if the values of the positive gap increase or the negative gap values decrease.

The independence-frequency gaps were first derived for 4 severity levels by 4 settings to explore the effects of severity and setting on the gaps. Then, for each severity level, the gaps were analyzed for diagnostic groups by settings to explore the effects of diagnosis and setting on the gaps. Distribution of the severity levels of the majority of the participants (n = 13,906, 76.7%) classified on the basis of the four major diagnostic groups is listed in Table 1. The mean gaps (positive gaps are shown with bold characters and numbers in Table 2) were analyzed across diagnosis/severity with ANOVA, and across setting with repeated measures ANOVA, followed by Scheffé’s post hoc analyses (Table 2). Considering the significant discrepancy of sample sizes between diagnostic/severity groups that could contribute to variance inequality, Leven’s test for homogeneity was conducted before ANOVA. Welch ANOVA and Games-Howell post hoc analyses were performed if the data failed to meet the equal variance assumption. The relationship between independence and frequency were analyzed with Pearson’s correlation in four settings for each severity level with alpha set at 0.05 (2-tailed).

In Step 3, only the children with mild severity (n = 7,719) were included to examine the nature of the independence-frequency gap as a function of the age by diagnosis/setting interaction with linear trend analysis. Trends were plotted by age on dual Y coordinates, contrasting the discrepancy between independence and frequency (Fig 3). Linear trends analyses with the significance level set at 0.05 (2-tailed) were conducted for independence, frequency, and the gaps across ages (Table 3). The significance of the linear trend was tested by ANOVA for 12 age groups, and the between-groups sum of squares for the effect of age was partitioned into a linear trend and higher order trends. The linear trend component was tested by an F-ratio (the mean square for linear trend/error term). The statistical analyses in Step 2 and Step 3 were performed with SPSS 20.0 (IBM SPSS, Chicago, Illinois, 2013).

**Results**

For Step 1, among the 18,119 children, 1,106 (6%) children had two of the four diagnoses, and only 16 (< 0.1%) children had more than two diagnoses. Among the children diagnosed with an intellectual disability (n = 10,310) (Table 1), 650 children (6%) were also diagnosed with ASD, 264 (3%) with language/hearing impairments, and 154 (1%) children with cerebral palsy. Furthermore, there was significant association between impairment severity and the number of
**Step 1: all enrolled children**  
(n= 18,119)  
Mean scores for capability and frequency were plotted on dual Y coordinates grouped by severity and settings across age (Figure 2)

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**Step 2: children having only one type of diagnosis**  
(n= 13,906)  
Capability-frequency gaps were derived for diagnostic group by settings for 4 severity levels of impairment (Table 2)

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**Step 3: children with mild severity of impairment and having only one type of diagnosis**  
(n= 7,719)  
Linear trends analyses across age grouped by diagnosis and settings (Table 3, Figure 3)
comorbid conditions (Chi-square = 1145.38, P < 0.001). Of the 9,561 children with mild severity, 7,720 children (81%) had only one diagnosis, while of the 1,170 children with profound severity, 540 children (46%) had two or more diagnoses. Of the 18,119 children, 45% lived in the northern part of Taiwan, which has the highest level of urbanization, 21% in the central part, 27% in the southern part, with 7% in the western part and some remote islands, which have the lowest level of urbanization.

Fig 2 provides the complete picture of the settings, severity level of impairment and the independence-frequency gap by age. A negative gap was found for children with mild levels of impairments for all settings (Fig 2 top row). As the gaps increased with age, the significance of the trends were further examined in Step 3. For children with moderate to profound levels of impairment, the trends varied by setting. A positive gap was found even for children with severe to profound levels of impairments, especially for children aged 6–9 years in home and school settings (Fig 2 lower rows in home and school settings).

The analyses for Step 2 for those children with only one diagnosis revealed a similar pattern to the whole group; that is, the mean gaps were all negative for children with mild to moderate levels of impairment, shifting to positive for children with severe level of impairment in the school setting, and were all positive for children with profound level of impairment (Table 2). Comparing gaps among different severity groups with ANOVA, there were significant differences among the 4 groups (Table 2, first box). Largest positive gaps were found for children with profound level of severity in all the 4 settings with the widest negative gaps for the mild group. Comparing gaps among different diagnostic groups, wider negative gaps were found among children with mild or moderate ID in the home, neighborhood/community, and school.
Table 2. Comparisons of the independence-frequency gaps across setting, severity and diagnosis for children with only one diagnosis of all severity levels in body function.

| Settingsa | Home | Neighborhood and community (NC) | Schoolb | HCLA | Significancec | Post hocd |
|-----------|------|--------------------------------|---------|------|--------------|-----------|
| Severity levels (n = 13906) | | | | | | |
| Mild (n = 7719) | -8.1 (± 14.93) | -16.6 (± 22.19) | -8.5 (± 17.35) | -8.2 (± 17.85) | F = 672.61, P < 0.001 | Home = HCLA = School > NC |
| Moderate (n = 4430) | -5.7 (± 15.39) | -12.1 (± 20.98) | -4.5 (± 18.13) | -4.4 (± 17.4) | F = 303.17, P < 0.001 | HCLA = School > Home > NC |
| Severe (n = 1129) | -1.0 (± 15.12) | -4.9 (± 18.98) | 2.2 (± 17.46) | -0.2 (± 16.01) | F = 51.21, P < 0.001 | School > HCLA = Home > NC |
| Profound (n = 628) | 3.3 (± 13.61) | 1.3 (± 14.88) | 6.6 (± 17.95) | 2.5 (± 12.39) | F = 15.51, P < 0.001 | School > Home = HCLA = NC |
| Significance | F = 168.70, P < 0.001f | F = 318.31, P < 0.001f | F = 232.53, P < 0.001f | F = 186.87, P < 0.001f | | |
| Post hoc | P > S > Mod > Milh | P > S > Mod > Milh | P > S > Mod > Milh | | | |
| Mild (n = 7719) | | | | | | |
| ID (n = 4966) | -8.7 (± 14.89) | -17.6 (± 22.04) | -8.9 (± 17.37) | -8.3 (± 17.52) | F = 499.75, P < 0.001 | HCLA = School = Home > NC |
| ASD (n = 2117) | -6.9 (± 14.50) | -15.1 (± 22.11) | -8.2 (± 17.02) | -7.8 (± 18.14) | F = 142.30, P < 0.001 | Home = HCLA > School > NC |
| LH (n = 531) | -7.1 (± 16.07) | -14.0 (± 23.24) | -6.2 (± 17.84) | -7.9 (± 19.05) | F = 31.42, P < 0.001 | School = Home = HCLA > NC |
| CP (n = 105) | -4.3 (± 16.88) | -15.2 (± 22.18) | -6.0 (± 16.54) | -5.1 (± 20.04) | F = 14.48, P < 0.001 | Home = HCLA = School > NC |
| Significance | F = 10.48, P < 0.001f | F = 9.88, P < 0.001f | F = 4.44, P = 0.004f | F = 1.67, P = 0.17f | | |
| Post hoc | CP = ASD = LH > IDh | LH = ASD = CP > IDg | CP = LH > ASD = IDg | CP = ASD = LH = IDg | | |
| Moderate (n = 4430) | | | | | | |
| ID (n = 3295) | -6.1 (± 15.37) | -13.0 (± 21.03) | -4.7 (± 18.26) | -4.4 (± 16.82) | F = 271.36, P < 0.001 | HCLA = School > Home > NC |
| ASD (n = 805) | -4.6 (± 15.20) | -8.2 (± 19.93) | -4.2 (± 16.80) | -3.4 (± 19.05) | F = 20.62, P < 0.001 | HCLA = School > Home > NC |
| LH (n = 205) | -4.8 (± 16.27) | -12.1 (± 21.78) | -4.2 (± 19.01) | -7.1 (± 18.68) | F = 14.23, P < 0.001 | School = Home = HCLA > NC |
| CP (n = 125) | -2.6 (± 15.05) | -11.4 (± 22.16) | -1.9 (± 21.02) | -4.1 (± 18.98) | F = 12.79, P < 0.001 | School = Home = HCLA > NC |
| Significance | F = 3.84, P = 0.01f | F = 12.22, P < 0.001f | F = 0.80, P = 0.49f | F = 2.14, P = 0.95f | | |
| Post hoc | CP = ASD = LH > IDh | ASD > CP = LH = IDh | CP = ASD = LH = IDh | ASD = CP = ID = LHh | | |
| Severe (n = 1129) | | | | | | |
| ID (n = 673) | -1.7 (± 16.12) | -5.3 (± 19.70) | 1.7 (± 17.93) | -0.5 (± 16.64) | F = 29.46, P < 0.001 | School > HCLA = Home > NC |
| ASD (n = 260) | -0.2 (± 13.16) | -3.7 (± 16.67) | 2.1 (± 16.83) | 1.5 (± 15.43) | F = 9.48, P < 0.001 | School = HCLA > Home > NC |
| LH (n = 124) | -1.0 (± 13.44) | -7.2 (± 21.98) | 2.7 (± 15.44) | -2.2 (± 16.01) | F = 9.15, P < 0.001 | School > Home = HCLA > NC |
| CP (n = 72) | 2.6 (± 14.44) | -1.4 (± 13.16) | 6.2 (± 18.56) | -2.1 (± 10.50) | F = 5.14, P = 0.002 | School = Home > NC = HCLA |

(Continued)
settings, and children with mild ASD in the school setting (Table 2, 2nd and 3rd box). A significantly greater positive gap was found only for the profound children with CP in home setting (Table 2, 5th box).

Across the four settings, the widest negative gap was found significantly in the setting of neighborhood/community for the four diagnostic groups of children with mild to severe level of impairments based on the post hoc analysis (Table 2). For children with a profound level of impairment, the largest positive gaps were in the school setting, especially for children with ID and ASD (Table 2, 5th box).

The correlations between independence and performance for the four settings were moderate ($r = 0.59–0.70$, $P < 0.001$) for children with mild severity, were moderate to high ($r = 0.66–0.76$, $P < 0.001$) for children with moderate severity, were high for both severe ($r = 0.77–0.84$, $P < 0.001$) and profound groups ($r = 0.75–0.84$, $P < 0.001$).

The results of the Step 3 analysis are shown in Table 3, with mean values of frequency, independence, gap and the corresponding trends with age for children with mild severity and

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### Table 2. (Continued)

| Settings $^a$ | Home | Neighborhood and community (NC) | School $^b$ | HCLA | Significance $^c$ | Post hoc $^d$ |
|--------------|------|--------------------------------|------------|------|------------------|--------------|
| Significance | F = 2.25, $P = 0.08^h$ | F = 2.47, $P = 0.06^i$ | F = 1.38, $P = 0.25^e$ | F = 2.29, $P = 0.77^g$ |
| Post hoc     | CP = ASD = ID = LH $^h$ | CP = ASD = ID = LH $^h$ | ID = ASD = LH = CP $^j$ | ASD = ID = CP |
| Profound (n = 628) | |
| ID (n = 318) | 3.0 (± 14.22) | 0.9 (± 15.89) | 6.0 (± 18.53) | 2.2 (± 12.46) |
| ASD (n = 149) | 1.9 (± 11.58) | 0.4 (± 13.10) | 4.9 (± 14.17) | 3.7 (± 13.42) |
| LH (n = 50) | 2.0 (± 13.90) | 2.7 (± 14.36) | 8.5 (± 20.36) | 1.0 (± 8.24) |
| CP (n = 111) | 6.7 (± 13.85) | 3.3 (± 14.28) | 10.6 (± 20.00) | 2.24 (± 12.16) |

$^a$ Values are mean (±SD) of the gap of the scores between independence and frequency, and the values in bold present positive gaps.

$^b$ The domains of school setting are only applicable for children with the occupation of student, therefore the number of participants (n = 13,520) of this domains would be less than the other domains (n = 13,906).

$^c$ The values of the gaps were compared across setting using repeated measured ANOVA.

$^d$ The post hoc analyses across settings were performed using Scheffé’s method.

$^e$ The values of the gaps were compared across diagnostic/severity groups using one-way ANOVA.

$^f$ The values of the gaps were compared across diagnostic/severity groups using Welch ANOVA because the data fail to meet the equal variance assumption by significant Levene’s test for homogeneity of variances.

$^g$ The post hoc analyses across diagnostic groups were performed across diagnostic groups with Scheffé’s method.

$^h$ The post hoc analyses across diagnostic/severity groups were performed using Games-Howell post hoc analysis because the data fail to meet the equal variance assumption by significant Levene’s test for homogeneity of variances.

HCLA = home and community living activities; ID = Intellectual disability; ASD = Autistic spectrum disorder; LH = language/hearing impairment; CP = Cerebral palsy; Mil = mild; Mod = moderate; S = severe; P = profound

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A decreasing trend of independence restriction with age was found for all diagnostic groups except children with CP at home and school settings, indicating that parents perceived improvement of the children’s ability in daily activities. The trends of frequency, however, are more variable than trends for independence across age groups. A review of the trend lines in Table 3 and Fig 3 indicated that the gaps for each setting by diagnostic group were significantly wider with age except for two diagnostic groups by settings (LH in school setting, and CP in HCLA setting).

**Discussion**

The results of this large-sample study provide a picture of independence and frequency of participation using a broad distribution of severity and diagnoses of childhood disabilities in Taiwan. About half of the sample represented children with ID (57%) or were mildly impaired in body functions/structures (53%).

For the analysis of Step 2 in this study, widest negative gaps between independence and frequency (indicating potentially more environmental barriers) were found for neighborhood/community settings than the other three settings for children with mild to severe levels of impairment (Table 2, first box), especially for children aged 6–9 years (Fig 2). School and home may be the settings that provide relatively sufficient support for the less severe and younger child to engage in activities. For children with profound levels of impairment, the discrepancy between independence and frequency even changed to positive gaps (indicating hypothesized environmental support) especially in school settings. It appears that children with severe and
Table 3. Mean scores of frequency, independence, and gaps between independence and frequency across setting and diagnosis and the trend analyses for children with mild severity in body function.

| Setting                           | Frequency Mean (±SD) | Trend with age | Independence Mean (±SD) | Trend with age | Independence-frequency gap a Mean (±SD) | Trend with age |
|-----------------------------------|----------------------|----------------|-------------------------|----------------|----------------------------------------|----------------|
| **Home**                          |                      |                |                         |                |                                        |                |
| ID                                | 30.8 (± 17.58)       | Up, F = 5.10, P = 0.02 | 22.1 (± 16.59) | Dn, F = 190.24, P < 0.001 | -8.7 (± 14.89) | Wider, F = 155.56, P < 0.001 |
| ASD                               | 34.3 (± 16.66)       | Up, F = 5.10, P = 0.02 | 27.4 (± 16.52) | Dn, F = 30.41, P < 0.001 | -6.9 (± 14.50) | Wider, F = 80.53, P < 0.001 |
| LH                                | 26.0 (± 17.21)       | Dn, F = 8.10, P = 0.01 | 18.9 (± 16.56) | Dn, F = 43.11, P < 0.001 | -7.1 (± 16.07) | Wider, F = 11.55, P = 0.001 |
| CP                                | 23.2 (± 18.57)       | n.s., F = 0.03, P = 0.87 | 18.9 (± 17.39) | n.s., F = 2.77, P = 0.10 | -4.3 (± 16.88) | Wider, F = 4.19, P = 0.04 |
| **Neighborhood and community**    |                      |                |                         |                |                                        |                |
| ID                                | 45.4 (± 24.04)       | Up, F = 4.44, P = 0.04 | 27.7 (± 23.3)  | Dn, F = 72.75, P < 0.001 | -17.6 (± 22.04) | Wider, F = 125.91, P < 0.001 |
| ASD                               | 52.8 (± 23.08)       | Up, F = 35.93, P < 0.001 | 37.8 (± 25.23) | Dn, F = 13.48, P < 0.001 | -15.1 (± 22.11) | Wider, F = 116.50, P < 0.001 |
| LH                                | 37.8 (± 26.08)       | n.s., F = 0.82, P = 0.37 | 23.7 (± 22.7)  | Dn, F = 20.97, P < 0.001 | -14.0 (± 23.24) | Wider, F = 12.20, P = 0.001 |
| CP                                | 36.5 (± 27.25)       | n.s., F = 0.53, P = 0.47 | 21.2 (± 21.7)  | Dn, F = 20.97, P < 0.001 | -15.2 (± 22.18) | Wider, F = 5.42, P = 0.02 |
| **School**                        |                      |                |                         |                |                                        |                |
| ID                                | 34.9 (± 21.03)       | n.s., F = 0.91, P = 0.34 | 26.0 (± 19.36) | Dn, F = 162.63, P < 0.001 | -8.9 (± 17.37) | Wider, F = 169.50, P < 0.001 |
| ASD                               | 37.2 (± 20.89)       | Up, F = 23.92, P < 0.001 | 29.0 (± 19.25) | Dn, F = 9.43, P = 0.002  | -8.2 (± 17.02) | Wider, F = 92.67, P < 0.001 |
| LH                                | 26.9 (± 21.63)       | n.s., F = 2.47, P = 0.12 | 20.7 (± 18.98) | Dn, F = 13.31, P < 0.001 | -6.2 (± 17.84) | n.s., F = 3.76, P = 0.05 |
| CP                                | 26.0 (± 23.50)       | n.s., F = 1.35, P = 0.25 | 20.0 (± 20.49) | n.s., F = 0.33, P = 0.57 | -6.0 (± 16.54) | Wider, F = 5.32, P = 0.02 |
| **Home and community living activities** |                |                |                         |                |                                        |                |
| ID                                | 46.0 (± 22.28)       | Dn, F = 437.71, P < 0.001 | 37.7 (± 23.23) | Dn, F = 890.45, P < 0.001 | -8.3 (± 17.52) | Wider, F = 118.65, P < 0.001 |
| ASD                               | 45.7 (± 22.10)       | Dn, F = 133.44, P < 0.001 | 37.9 (± 22.24) | Dn, F = 330.79, P < 0.001 | -7.8 (± 18.14) | Wider, F = 50.81, P < 0.001 |
| LH                                | 40.6 (± 23.98)       | Dn, F = 64.85, P < 0.001 | 32.7 (± 24.56) | Dn, F = 131.65, P < 0.001 | -7.9 (± 19.05) | Wider, F = 10.51, P = 0.001 |
| CP                                | 41.6 (± 25.11)       | n.s., F = 1.16, P = 0.28 | 36.5 (± 25.26) | Dn, F = 5.85, P = 0.018  | -5.1 (± 20.04) | n.s., F = 3.04, P = 0.09 |

a Values are showed for the scores of independence minus the scores of frequency

Up = significantly upward trend; Dn = significantly downward trend; ns = non-significant; ID = Intellectual disability (n = 4966); ASD = Autistic spectrum disorders (n = 2117); LH = language/hearing impairment (n = 531); CP = Cerebral palsy (n = 105).

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Profound disabilities receive more support in school settings (Table 2, 1st and 4th-5th boxes). Previous studies have shown that children with disabilities experience barriers for participation in different settings, especially with regard to physical demands and needs in school. These areas were largely unmet in accommodations of special computer equipment, adapted furniture or accessibility of school buildings [24,42–44]. In this study, however, the children with ID or ASD with mild severity were even less supported than children with physical disability in social participation (Table 2, 2nd boxes).

Social and institutional barriers have been identified as the most significant environmental barriers in studies of children with disabling conditions in Canadian communities [26,45]. Parents reported that their children encountered greater barriers to participate in school and work environments, and the barriers included problems with service, assistance, attitude, and policy. Poorly designed barrier-free community facilities, difficulty getting information about services and programs, the vague process of applying to schools, and negative attitudes were participation challenges faced by children in these studies [26,45]. The attitudes and abilities of teachers to modify courses and instruction were also important influential factors on the extent of children’s participation [26,45].

Further statistical analyses in Step 3 indicated that the independence restriction to participate in daily activities decreased significantly with age among the children with mild severity, except for children with CP in home and school settings. The trend of frequency with age, however,
varied across the impairment by diagnosis clusters (Table 3). The trend of gaps showed that the mean frequency restriction of participation was higher than mean independence across settings and diagnoses and the mean independence-frequency gap became progressively wider with age across settings (Fig 3). This may suggest that children with mild severity have fewer opportunities to do tasks than they are capable of doing as they transition to adulthood.

It should be noted that for children with mild severity of impairment, data for independence and frequency for children with ID and ASD were relatively more restricted than for children with CP or LH (Table 3). The negative gap was also more obvious in children with mild rather than severe impairment (Table 2). This may be due to the social nature of participation and the fact that the children with ID or with mild impairment face more environmental barriers (suggested by negative gaps) than other diagnoses or more severe groups.

In the past, eligibility for financial or social support in Taiwan was based on either impairment or diagnosis, and the children with more severe levels of impairment or with more observable diagnoses such as motoric (e.g., cerebral palsy) or sensory deficits (e.g., hearing loss or visual impairments) were eligible for packaged services (especially pension or assistive technology or home services) matched to specific diagnoses and impairment levels.

Resource distribution, however, should be based on the consideration that children with social/emotional as well as less severe impairments also need tailored services (such as personal support and care)[11] aimed at the goal of optimal participation as framed by ICF-CY and consistent with age-appropriate milestones. The joint impairment-participation decision for the Disability Evaluation System, particularly for the large population of individuals with mild intellectual disability is now under construction in Taiwan [18]. The results of this study provide evidence to inform policy and decision making for the DES.

Overall, based on the profiles of independence and frequency among children with disabilities provided by Step 1 (Fig 2), restriction of independence in daily participation decreased with age whereas restriction of participation frequency varied across setting and diagnosis. These data suggest that frequency, defined as the performance dimension of participation, what children did in the past 6 months, is a more useful variable and context-dependent measure of participation than independence. The negative gap between independence-frequency became wider with age, especially in neighborhood and community settings and in children with mild impairment. Children with mild levels of impairments are probably more often in neighborhood and community settings and thus exposed to more complex and less adapted environments. A positive gap for participation was found for children with severe to profound levels of impairment in home and school settings especially children aged 6–9 years of age. It may be that families and schools provide more support to children of a younger age and with more severe impairment levels.

A group of children sharing the same diagnosis, such as CP, may show a wide range of disability/functioning in daily life and some may not have limitations in mobility. Children with various diagnoses may also share common limitations of functioning, especially for participation in specific settings because of the environmental support or barriers encountered.

Children’s participation provides valuable information beyond diagnoses for assessing children’s needs [46] in all education settings (classroom settings as well as school-related clinical settings) with implications for continuity in the transitions from one educational level to the next and into work and employment [47]. Further analysis of environmental factors other than physical settings is needed, such as parents’ perspective on the provision of opportunities for activities. Continued research on the social as well as physical factors of the environment is needed to identify the conditions and opportunities that can promote the participation of children with disabilities.

Several limitations of this study need to be considered. First, the distribution of children across levels of severity, diagnoses, and ages was not even. This resulted in varied mean scores...
as shown in Fig 3 especially for the children with CP and in a specific age range. Second, the comorbidity issue is an inevitable challenge in data analysis for the effects of diagnoses on functional outcomes. Therefore, we only performed statistical analysis for Step 2 and 3 where children with more than one type of diagnosis were excluded. Third, the FUNDES-Child was designed to obtain caregiver’s or parents’ perceptions of children’s independence and frequency of engaging in activities. Thus, all the information was based on ratings by caregivers or parents within the scope of the report format of the FUNDES-Child in Taiwan. Caregivers or parents may not be as familiar with children’s participation in the school setting as in the home. Self-rating of participation by children was not feasible in that children often would have difficulty comprehending the questions.

Parents rating for measuring children’s participation has been seen as a valid approach to explore the patterns of participation in different settings and ages [48,49]. For children capable of answering survey questions, they should respond for themselves [9] as the integration of the child’s perspective and choice is especially important for full participation in life as an active learner [32,50]. Future studies of FUNDES-Child will prioritize self-ratings of participation by children, perhaps using pictures of the activities. Fourth, the independence-frequency gap may be only one of several ways to conceptualize the role of the environment. While we believe this is one reasoned approach, others may emerge as these concepts develop over time. The relations between gaps and environmental factors need further studies.

Conclusion

In conclusion, this large-sample study of participants drawn from a disability register system encompassing a broad range of disabilities, demographic characteristics, and ages provides a rather comprehensive picture of functional characteristics/limitations of children with developmental disabilities in Taiwan. Diagnoses, physical conditions and related severity of impairments and limitations of functioning and participation have been differentiated with the ICF/ICF-CY model [4,21] and validated with empirical data [5,17]. Findings for this study indicate that children’s participation depends not only on the severity of their impairments or diagnoses, but also on their age, the settings and supports provided by their environments. A priority for future research is thus further identification of environmental opportunities that positively impact frequency of participation in daily activities for children with developmental disabilities.

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Author Contributions

Conceived and designed the experiments: AWH HFL THL SWT WTC. Performed the experiments: AWH HFL THL SWT WTC CFY WCC LJJK TFW. Analyzed the data: AWH CFY WCC. Contributed reagents/materials/analysis tools: HFL AWH LJJK CFY TFW. Wrote the paper: AWH HFL RJS DJL CFY LJJK. Research project organization: HFL THL SWT WTC. Research project execution: HFL THL AWH.

References

1. Winter S, Autry A, Boyle C, Yeargin-Allsopp M (2002) Trends in the prevalence of cerebral palsy in a population-based study. Pediatrics 110: 1220–1225. PMID: 12456922
2. Crocker AC, Rubin IL (1989) The spectrum of medical care for developmental disabilities. In: Crocker AC, editor. Developmental disabilities: Delivery of medical care for children and adult. Philadelphia, PA: Lea and Febiger. pp. 10.

3. Simeonsson RJ, Carlson D, Huntington GS, McMillen JS, Brent JL (2001) Students with disabilities: A national survey of participation in school activities. Disabil Rehabil 23: 49–63. PMID:11214716

4. World Health Organization (2007) International Classification of Functioning, Disability and Health: Children & Youth Version (ICF-CY). Geneva: WHO.

5. McDougall J, King G, de Wit DJ, Miller LT, Hong S, Offord DR, et al. (2004) Chronic physical health conditions and disability among Canadian school-aged children: A national profile. Disabil Rehabil 26: 35–45. PMID:14660197

6. Lollar DJ, Hartzell MS, Evans MA (2012) Functional difficulties and health conditions among children with special health needs. Pediatrics 129: e714–e722. doi: 10.1542/peds.2011-0780 PMID:22371461

7. Boyle CA, Boulet S, Schieve LA, Cohen RA, Blumberg SJ, Yeargin-Allsopp M, et al. (2011) Trends in the prevalence of developmental disabilities in US children, 1997–2008. Pediatrics 127: 1034–1042. doi: 10.1542/peds.2010-2989 PMID:21606152

8. Boyle CA, Decoufle P, Yeargin-Allsopp M (1994) Prevalence and health impact of developmental disabilities in US children. Pediatrics 93: 399–403. PMID:7509480

9. Newacheck PW (1998) Prevalence and impact of disabling chronic conditions in childhood. AJPH 88: 610–617.

10. Ministry of Health and Welfare (2007) People with Disabilities Rights of Protection Act. Taiwan. Available: http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?PCODE=D0050046. Accessed 30 July 2014.

11. Chiu WT, Yen CF, Teng SW, Liao HF, Chang KH, Chi WC, et al. (2013) Implementing disability evaluation and welfare services based on the framework of the international classification of functioning, disability and health: Experiences in Taiwan. BMC Health Serv Res 13: 416. doi: 10.1186/1472-6963-13-416 PMID:24125482

12. Hwang AW, Liou TH, Bedell GM, Kang LJ, Chen WC, Yen CF, et al. (2013) Psychometric properties of the Child and Adolescent Scale of Participation—Traditional Chinese Version. Int J Rehabil Res 36: 211–220. doi: 10.1097/MRR.0b013e32835d0b27 PMID:23328815

13. Liao HF, Yen CF, Hwang AW, Liou TH, Chang BS, Wu TF, et al. (2013) Introduction to the application of the Functioning Scale of the Disability Evaluation System. Formosan J Med 17: 317–331.

14. Bedell G (2004) Developing a follow-up survey focused on participation of children and youth with acquired brain injuries after discharge from inpatient rehabilitation. NeuroRehabilitation 19: 191–205. PMID:15502253

15. Bedell G (2009) Further validation of the Child and Adolescent Scale of Participation (CASP). Dev Neurorehabil 12: 342–351. doi: 10.3109/17518420903087277 PMID:20477563

16. Ministry of Health and Welfare (2014) Disability statistics. Taiwan. Available: http://www.mohw.gov.tw/cht/DOS/Statistic.aspx?f_list_no=312&fod_list_no=4198. Accessed 30 July 2014.

17. Van Naarden Braun K, Yeargin-Allsopp M, Lollar D (2009) Activity limitations among young adults with developmental disabilities: A population-based follow-up study. Res Dev Disabil 30: 179–191. doi: 10.1016/j.ridd.2008.02.004 PMID:18455365

18. Chi WC, Liou TH, Huang WNW, Yen CF, Teng SW, Chang IC (2013) Developing a disability determination model using a decision support system in Taiwan: A pilot study. J Formos Med Assoc 112: 479–481. doi: 10.1016/j.jfma.2013.06.005 PMID:23871551

19. Law M, Finkelman S, Hurley P, Rosenbaum P, King S, King G, et al. (2004) Participation of children with physical disabilities: Relationship with diagnosis, physical function, and demographic variables. Scand J Occup Ther 11: 156–162.

20. United Nations (1989) Convention on the Rights of the Child (UNCRC) (Article 7, 8, 12, 13, 17, 23).

21. World Health Organization (2001) International Classification of Functioning, Disability and Health: ICF. Geneva: WHO.

22. Bedell G, Dumas HM (2004) Social participation of children and youth with acquired brain injuries discharged from inpatient rehabilitation: A follow-up study. Brain Inj 18: 65–82. PMID:14660237

23. Eriksson L (2005) The relationship between school environment and participation for students with disabilities. Pediatr Rehabil 8: 130–139. PMID:16089253

24. Galvin J, Froude EH, McAleer J (2010) Children's participation in home, school and community life after acquired brain injury. Aust Occup Ther J 57: 118–126. doi: 10.1111/j.1440-1630.2009.00822.x PMID:20854577
25. Kang LJ, Palisano RJ, Orlin MN, Chiarello LA, King GA, Polansky M (2010) Determinants of social participation with friends and others who are not family members for youths with cerebral palsy. Phys Ther 90: 1743–1757. doi: 10.2522/ptj.20100048 PMID: 20930051

26. Law M, Haigh M, Milroy B, Willms D, Stewart D, Rosenbaum P (1999) Environmental factors affecting the occupations of children with physical disabilities. J Occup Sci 6: 102–110.

27. Bedell G (2013) Community participation, supports, and barriers of school-age children with and without disabilities. Arch Phys Med Rehabil 94: 315–323. doi: 10.1016/j.apmr.2012.09.024 PMID: 23044364

28. Coster W, Law M, Bedell G, Liljenquist K, Kao YC, Khetani M, et al. (2013) School participation, supports and barriers of students with and without disabilities. Child Care Health Dev 39: 535–543. doi: 10.1111/chc.12046 PMID: 23763254

29. Hwang AW, Granlund M, Liao HF, Kang LJ (2013) Measures of “Participation” in school children with disabilities. FJPT 38: 37–49.

30. Whiteneck G, Dijkers MP (2009) Difficult to measure constructs: Conceptual and methodological issues concerning participation and environmental factors. Arch Phys Med Rehabil 90: S22–35. doi: 10.1016/j.apmr.2009.06.009 PMID: 19892071

31. Holsbeeke L, Ketelaar M, Schoemaker MM, Gorter JW (2009) Capacity, capability, and performance: Different constructs or three of a kind? Arch Phys Med Rehabil 90: 849–855. doi: 10.1016/j.apmr.2008.11.015 PMID: 19406307

32. Morris C (2009) Measuring participation in childhood disability: How does the capability approach improve our understanding? Dev Med Child Neurol 51: 92–94. doi: 10.1111/j.1469-8749.2009.03461.x PMID: 19740215

33. Palisano RJ, Chiarello LA, Orlin M, Oeffinger D, Polansky M, Maggs J, et al. (2010) Determinants of intensity of participation in leisure and recreational activities by children with cerebral palsy. Dev Med Child Neurol 53: 142–149. doi: 10.1111/j.1469-8749.2010.03819.x PMID: 20964676

34. Palisano RJ, Orlin M, Chiarello LA, Oeffinger D, Polansky M, Maggs J, et al. (2011) Determinants of intensity of participation in leisure and recreational activities by youth with cerebral palsy. Arch Phys Med Rehabil 92: 1468–1476. doi: 10.1016/j.apmr.2011.04.007 PMID: 21878219

35. Fauconnier J, Dickinson HO, Beckung E, Marcelli M, McManus V, Michelsen SI, et al. (2009) Participation in life situations of 8–12 year old children with cerebral palsy: Cross sectional European study. BMJ (Online) 338: 1116–1121.

36. Tucker CA, Cieza A, Riley AW, Stucki G, Lai JS, Bedirhan Ustun T, et al. (2014) Concept analysis of the Patient Reported Outcomes Measurement Information System (PROMIS) and the International Classification of Functioning, Disability and Health. Qual Life Res 23: 1677–1686. doi: 10.1007/s11136-014-0622-y PMID: 24500657

37. Teng SW, Yen CF, Liao HF, Chang KH, Chi WC, Wang TH, et al. (2013) Evolution of system for disability assessment based on the International Classification of Functioning, Disability, and Health: A Taiwanese study. J Formos Med Assoc 112: 691–698. doi: 10.1016/j.jfma.2013.09.007 PMID: 24099681

38. Ministry of Health and Welfare The classification, categories, grades of disability and standards in disability evaluation system. Taiwan. Available: http://law.moj.gov.tw/Law/LawSearchResult.aspx?p=A&t=A1A2E1F1&k1=%E9%91%91%E5%AE%9A%E5%9D%91%E5%BA%A6. Accessed 30 July 2014.

39. Bedell G (2011) The Child and Family Follow-up Survey (CFFS)-Administration and Scoring Guidelines [Unpublished Manual].

40. Liao HF, Fan CJ, Liou TH, Yen CF, Wu TF, Chang BS, et al. (2013) Training programs for testers of the Functioning Scale of the Disability Evaluation System in Taiwan and their outcomes. Formosan J Med 17: 368–380.

41. Palisano RJ, Tieman BL, Walter SD, Bartlett DJ, Rosenbaum PL, Russell D, et al. (2003) Effect of environmental setting on mobility methods of children with cerebral palsy. Dev Med Child Neurol 45: 113–120. PMID: 12578237

42. Tieman B, Palisano RJ, Rosenbaum P (2003) The impact of environmental setting on the mobility of children with cerebral palsy: Research findings and clinical implications. CanChild Centre for Childhood Disability Research.

43. Imms C (2008) Children with cerebral palsy participate: A review of the literature. Disabil Rehabil 30: 1867–1884. doi: 10.1080/09638280701673542 PMID: 19037780
45. Law M, Petrenchik T, King G, Hurley P (2007) Perceived environmental barriers to recreational, community, and school participation for children and youth with physical disabilities. Arch Phys Med Rehabil 88: 1636–1642. PMID: 18047879

46. Almqvist L, Granlund M (2005) Participation in school environment of children and youth with disabilities: A person-oriented approach. Scand J Psychol 46: 305–314. PMID: 15842421

47. World Health Organization (2013) How to use the ICF: A practical manual for using the International Classification of Functioning, Disability and Health (ICF). Geneva: WHO.

48. McDougall J, Bedell G, Wright V (2013) The youth report version of the Child and Adolescent Scale of Participation (CASP): Assessment of psychometric properties and comparison with parent report. Child Care Health Dev 39: 512–522. doi: 10.1111/cch.12050 PMID: 23763252

49. Scorza P, Stevenson A, Canino G, Mushashi C, Kanyanganzi F, Munyanah M, et al. (2013) Validation of the "World Health Organization Disability Assessment Schedule for Children, WHODAS-Child" in Rwanda. PLoS One 8: e57725. doi: 10.1371/journal.pone.0057725 PMID: 23505437

50. Morris C (2007) Measuring children’s participation. Dev Med Child Neurol 49: 645–645. PMID: 17718817