ABSTRACT
Children with developmental disabilities may need extra care due to delayed development of self-care skills and/or a need for special care. Delayed self-care skills mean that care needs that are expected to diminish in typically developing children, persist.

Whether a child is independent in an aspect of self-care would be dependent on his chronological and developmental age, developmental disability and the associated impairments, as well as cultural and other environmental factors, such as the child's gender, the presence of a domestic helper, early attendance at preschool. Family circumstances should also be taken into consideration.

The functional assessment of ADL in infants and children with developmental disabilities has been distilled into the domains of Washing/Bathing, Dressing, Feeding, Toileting, Transferring and Mobility. They reflect the activities that occur in the typical day in the life of a child. Common developmental disabilities and possible effects on acquisition of independent ADL are discussed.

Keywords: Assessment, activities of daily living (ADL), self-care, developmental disability, children.

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INTRODUCTION
All children have a right to care by a parent or trusted adult. As infants, this care extends to every aspect of the child's life: bodily functions, and physical, mental and emotional development. As the child grows, the care required changes and generally diminishes, so that the child needs less and less help from the adults around him. Children with developmental disabilities require the same care and parenting as other children, with the addition of extra care due to impairments or chronic ill health. The extra care required may arise from delayed development of self-care skills and/or a need for special care. Delayed self-care skills mean that care needs - which are expected to diminish in typically developing children - persist. Special care needs 'go beyond the bounds of ordinary parenting', for instance involving ventilators or feeding tubes.

Developmental disabilities are a group of chronic disorders of early onset estimated to affect 5 to 10% of children. In more technical terms, these children often have a variety of associated impairments that may sometimes lead to limitations in the capacities of the child. According to the International Classification of Functioning, Disability and Health (ICF), capacities describe the individual's ability to execute a task or an action, and are defined as the highest probable level of functioning that a person may reach in a given domain at a given moment in a 'standardised or assumed' environment. Limitations in the capacities of the child will lead to the need for additional care. This additional care has been defined as 'dependency on medication or special diet, medical technology, assistive devices, personal assistance, need for medical care or related services or educational services over and above the usual for the child's age, or for special ongoing treatments, interventions or accommodations at home or in school'.

In a study of the families of 40000 disabled children in the UK, the majority of the children required extra assistance or supervision with multiple areas of daily life. The study also indicated variety in extra care tasks undertaken (physical help, supervision, guidance) and causal factors (physical limitation, cognitive difficulties, behavioural problems). In particular, the findings indicated that parents want professionals to recognise and offer explicit acknowledgement of the extra care they give their disabled children.

Certainly, taking into consideration the child's family circumstances is important. Families often have more than one child, and sometimes, more than one disabled child. Single parents may often have to care for their disabled children alone. Besides recognising and acknowledging the care that parents provide for their children with disabilities, it behooves us to offer family-centred care and assist with decision-making in a family-centric and not merely child- or patient-centric way.

INDEPENDENCE IN ADL IN INFANCY AND CHILDHOOD
Whether a child is independent in an aspect of ADL would be dependent on his chronological and developmental age, as well as cultural and other environmental factors. In typically developing children, there is generally an age range at which a certain ADL is achieved, but again, external cultural and other environmental factors need to be taken into consideration. In a study on normal Hong Kong Chinese children, some gender and culturally specific aspects (e.g. the presence of a maid at home, early attendance at preschool) were found to impact on the performance of functional skills. In general, girls mastered self-care and communication competencies before boys.

Crucial to measuring functional limitations is defining
essential daily tasks. These may include self-care activities of feeding (using a cup, spoon, and fork), grooming (washing face and hands, brushing teeth, combing hair), dressing upper body (shirts/blouses), dressing lower body (underwear, pants, shoes and socks), adjusting clothes, maintaining bladder and bowel continence, and cleaning up after toileting. Motor activities include changing positions from chairs, getting on and off the toilet or potty, getting in and out of the bathtub or shower, walking indoors and outdoors, and ascending and descending stairs.

The list of these activities reflects that during a typical day, a child will:

• complete a meal independently using appropriate utensils or be assisted in feeding
• wash hands and face, brush teeth, comb hair, or be assisted in these areas
• will dress and maintain continence or require assistance or diapers
• generally ambulate indoors and outdoors, get in and out of chairs, on and off toilets, in and out of showers, negotiate stairs, or be assisted with these tasks

EXISTING INSTRUMENTS

Currently, pediatric health professionals have several choices for adaptive-functional instruments. These include the WeeFIM (Functional Independence Measure), the Pediatric Evaluation of Disability Inventory (PEDI), the Battelle Developmental Inventory (BDI), and the Vineland Adaptive Behavior Scales (VABS). The PEDI, BDI and VABS are maximum data sets and involve detailed and extensive queries of self-care, mobility, communication and social items. On average, they require >30 minutes of interview time and complex scoring5,6. The WeeFIM is useful in assessing functional independence in children aged 6 months to 7 years (up to 21 years in people with developmental disabilities), and can be administered by a trained interviewer in approximately 15 minutes7. A simple screening questionnaires, disabilities), and can be administered by a trained interviewer in

FUNCTIONAL ASSESSMENT

Washing/Bathing

Typically developing children in Singapore often start bathing independently from the age of six or seven. Children exposed to day care may bathe independently earlier.

Extra care with washing may be needed because of a physical inability to wash or extra care for delicate skin. Frequent vomiting, diarrhoea, or weeping wounds necessitate repeated washing.

Dressing

From about 18 months, toddlers often assist with dressing and undressing, taking off shoes. By the age of three, he is able to pull his pants down and up, but needs help with buttons and other fastenings. Most five to six-year-olds are able to undress and dress independently.

Extra help when dressing may include practical help because of stiff or floppy limbs, repeated changes due to incontinence or dribbling, or guidance for a visually impaired child.

Feeding

An 18-month-old toddler can hold a spoon and get food safely to his mouth, although he may also play with the food. He is able to hold his cup between both hands and drink without much spilling. By the age of two, he is able to feed himself competently with a spoon, but is easily distracted. He is able to lift a cup and drink well without spilling, and replace the cup on the table without difficulty. He is able to ask for food and drink. By the age of four, he is able to eat skillfully with a fork and spoon.

Locally, however, many children have the skills to feed themselves and will do so in their preschool or child care settings, but at home, they are often fed by an adult caregiver, as it takes less time and involves less cleaning up. Many children continue to have milk feeds from an infant milk bottle till the age of sometimes five to six years, though they will drink from a cup during the day.

A child who finds chewing and swallowing difficult or requires tube feeding needs extra care at mealtimes, while children with behavioural disorders or insatiable (e.g. in Prader Willi Syndrome), or inappropriate (e.g. Pica) appetites require close supervision.

Toileting

Many toddlers begin to give notice of urgent toilet needs by restlessness and vocalisation. Bowel control may be attained but this is variable. They may indicate wet or soiled pants.

Encopresis is defined as regular soiling of the underwear in children aged four years and older, with no organic disease. The soiling can range from staining the underwear to semi-formed or formed stools. Enuresis is defined as a lack of urinary continence beyond the age of four for diurnal enuresis, a lack of urinary continence beyond the age of six for nocturnal enuresis, or the loss of continence after at least three months of dryness.
Young disabled children may need extra care when toileting due to bladder and/or bowel incontinence, stoma care, catheterisation or poor skin. Although most children are toilet trained by the age of three, this process may be prolonged or unsuccessful for disabled children.

**Transferring and Mobility**

Toddlers from the age of about 18 months are generally able to walk well and run carefully, with rapid improvement of locomotor skills from about two and a half years. Two year olds enjoy climbing, and will climb forward into an adult’s chair, then turn round and sit. By the age of four, they are able to walk or run alone up and down stairs, one foot to a step in adult fashion.

Inaccessible toilet facilities may mean an older physically disabled child continues to require adult assistance, while other disabled children find it hard to manage personal hygiene alone.

**Additional Care**

For children with special needs may also include the following:
- Extra care at night, e.g. turning an immobile child, giving medication or tube feeds
- Supervision for children with special needs when they are awake, as some may have frequent seizures, or lack an awareness of danger
- Behavioural problems ranging from inconsolable and constant crying in younger children to self-harm, destructiveness and violent or attention seeking behavior in older children.

**Types of Disability in Children and Infants**

In the form for application for FDW levy concession for persons with disabilities, the disability types have been divided into six main categories.

**Physical**

These would include congenital structural malformations as well as neuromuscular disorders, which affect motor function. In general, these disorders present fairly early, e.g. structural malformations such as spina bifida and arthrogryposis multiplex congenital would be picked up in the neonatal period. Other ‘physical’ disabilities such as cerebral palsy, Duchenne Muscular Dystrophy and other neuromuscular disorders generally present in the first to third year of life.

**Intellectual**

There is some confusion about the label “Global Developmental Delay” (GDD). This is a subset of developmental disabilities defined as significant delay in two or more of the following developmental domains: gross/fine motor, speech/language, cognition, social/personal, and activities of daily living. GDD describes a clinical presentation that has a heterogeneous aetiologic profile and is associated with age-specific deficits in adaptation and learning skills. Those deficits are evident in comparison with the skills attainment of chronological peers. The term GDD is usually reserved for younger children (i.e. typically less than five years of age), whereas the term intellectual disability is usually applied to older children when IQ testing is more valid and reliable. However, a young child with the clinical picture of GDD is not necessarily destined to be intellectual disabled. Infants and children may have GDD owing to conditions such as cerebral palsy, certain neuromuscular disorders, and other conditions such as early environmental deprivation, yet when they are old enough for cognitive testing, they do not score within the range of intellectual disability.

**Autism Spectrum Disorder**

As this is a ‘spectrum disorder’, the symptoms, abilities, and characteristics are expressed in many different combinations and in any degree of severity. Hence, children with autism vary widely in their cognitive, language, and social abilities, with a variety of strengths and weaknesses. At one end of the spectrum, we may find a mute child, crouched in a corner of the room, spinning a paper clip over and over again for hours; at the other end of the spectrum is a highly verbal, mathematically and scientifically gifted boy in the Integrated Programme, but with significant social difficulties.

Their major deficit is in interactive social relationships and social communication, both verbal and non-verbal. Many children with ASD also have restricted interests and activities.

**Sensory**

This category is generally fairly straightforward, and would include children with a single sensory impairment, such as Visual or Hearing Impairment.

**Multiple Disabilities**

Children with multiple disabilities, such as coexisting cerebral palsy with mental retardation, or children with two sensory impairments, would fall into this category.

**Illustrations Using Case Studies**

**CASE 1. Spina Bifida**

V is a 7-year-old Indian girl with T12 spina bifida diagnosed only at delivery. This was surgically repaired on day 1 of life. A ventriculoperitoneal shunt was also inserted for hydrocephalus. She is presently attending primary one at a mainstream primary school, with therapy support from Teach Me services. Her father works shifts as a security guard whilst her mother is a homemaker, and her main caregiver. V has an older brother and a younger sister.

She is able to use a cup, spoon and fork without help; her mother cuts her meat. She independently brushes her hair,
CASE 2. Down Syndrome

AR is a four-year-old Malay boy with regular trisomy 21, delivered at term. He has a surgically repaired atrioventricular septal defect and is the fourth child of mature parents, who are very supportive and understand his disability.

Presently, AR is attending an EIPIC programme twice a week. He walked independently at age two, and has a steady gait. However, he is unaware of danger, and often climbs onto tables and chairs, to look out of the window. He is able to address his family members, and uses mainly single words, having difficulty with receptive and expressive language skills. His attention span is short, and he is unable to follow most instructions. His behaviour is often self-directed and he has difficulties with unsupervised play and problem solving. AR needs help with dressing, he is still on diapers throughout the day, and tries to use a spoon to feed himself, but is messy. Bath time is often prolonged and challenging, as AR often refuses to get out of the bath. AR’s mother has reported that it is a challenge looking after him, and that she is often exhausted at the end of the day.

Points to note:
- Cognitively high-functioning child, but needing help with many self-care activities.
- Supportive mother who is enabled, in an intact home environment.

CASE 3. Autistic Spectrum Disorder

A is a 13-year-old Chinese girl with Autistic Spectrum Disorder with moderate intellectual disability, diagnosed when she was 3 years old. Her father works long hours as a taxi driver. Her mother is a homemaker, and is A’s main caregiver. She has two other younger typically developing siblings.

A is attending classes in a special school. She takes the school bus, but needs her mother to wait with her at the bus stop, as she sometimes still steps onto the road without looking to see if cars are coming. She is able to indicate some of her basic needs by gesture or signing, but has no speech. She has learned to wash, dress, and feed herself. She has also learned to cut out shapes with a pair of scissors. However, she has not learned to read, write or speak. At school, she is said to be compliant, and able to follow the school routines and schedules well. Too well, sometimes, as A is upset if there are changes in routine.

When she turned 12, A developed epilepsy. This is fairly well controlled on anticonvulsant medication. However, she also began to have more frequent and severe temper outbursts. She would scream, shout, throw objects and even push her mother. Often, it would take more than an hour for A to calm down. It was often difficult for A’s mother to tend to her needs and to those of her siblings, as well as the household chores. A was finally brought to see the child psychiatrist, who prescribed medication that has helped her tantrums. A daily routine and structure were also established for A, with scheduled times for waking up, washing, meals and watching television.

Points to note:
- Although A is a teenager, she requires care that is usually extended to perhaps a four-year-old.
with plus disease, needing laser photocoagulation, and bilateral sensorineural hearing loss. He was finally discharged when he reached a weight of 2 kg, with home oxygen for chronic lung disease. JX was weaned off the oxygen when he was about eight months old. Cranial ultrasound showed bilateral cystic periventricular leucomalacia.

His parents separated a year after JX was discharged from hospital. He had multiple readmissions for exacerbations of his chronic lung disease. Presently, his mother is working part-time as a sales promoter, and his maternal grandmother takes care of him whilst his mother is at work. However, grandma is in her mid-seventies, and also has medical problems of her own. JX’s father does not contribute to or participate in his care.

JX is attending the Programme for Children with Multiple Handicaps at Rainbow Centre. He has minimal self-mobility, and spends most of his waking day sitting up in a wheelchair. He is totally dependent in all activities of daily living, and although he has no speech, he is socially responsive and recognises his caregivers.

Points to note:

- Multiple severe disabilities, long term prognosis quite clear - will require lifelong care for most areas of ADL
- Single parent, with little family or community support

CONCLUSION

Children with developmental disabilities are a heterogeneous group with diverse clinical presentations, abilities and difficulties. It is important to consider each child and family individually. Care should be taken to consider the impact of the child’s impairments on requirements for extra care across all areas of daily life, as well as the individual family circumstances.

LEARNING POINTS

- Children with developmental disabilities often show a variety of associated impairments that may result in a lifelong need for additional care.
- Parents want professionals to recognise and offer explicit acknowledgement of the extra care they give their disabled children.