PARENTS' PSYCHOSOCIAL PROBLEMS INFLUENCING THE CONTINUITY OF EARLY INTERVENTION TREATMENT IN CHILDREN WITH DEVELOPMENTAL DELAY

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Developmental delay is a chronic disorder, which has a significant and continuing impact on the developmental progress of children. Whether the developmentally delayed children have continuous early intervention treatment is important for the success of early intervention. Professionals in early intervention services have been encouraged to understand parents' psychosocial problems and to further enhance their understanding of care for their delayed children. The purposes of this study were two-fold. The first was to identify the parents' psychosocial problems. The second purpose was to find parents' psychosocial variables that could be used to significantly discriminate whether the retarded children have continuously received treatment programs. A questionnaire to investigate parents' psychosocial problems was implemented. It mainly consisted of twenty descriptive items concerning parents' psychosocial conditions. Parents used a four-point Likert scale to rate the likelihood of the items. Parents also responded as to whether their child had received treatments continuously. Subjects were the parents of children who were diagnosed to have developmental delay. Factor analysis and discriminant analysis was used for statistical data examination. Analyzed data were from 207 completely finished questionnaires. Six parents' psychosocial factors were extracted via factor analysis, and they accounted for 46.9% of the variance of the twenty items. The factors were identified as family income, family members' relationships, caring skills, supporting resources, acceptance of the retarded child, and self-psychological adaptation. About half of the children had received treatment continuously. Discriminant analysis revealed that family income was crucial to parents' capacity to meet the needs for treatment.

Key words: developmental delay, early intervention, parents' psychosocial problems

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Developmental delay comprises a heterogeneous group of related early-onset chronic disorders that share the essential feature of a predominant disturbance in the acquisition of cognitive, motor,

language, or social skills. Developmental delay has a significant and continuing impact on the developmental progress of children [1]. Long-term early intervention treatments always encompass therapy or educational programs which provide a range of stimulation and training activities for the developmentally delayed children aged under six, and for their families [1, 2].

The reported benefits of early intervention are that it can increase the developmental and educational gains for retarded children [3, 4] and also improve the functions of families [5, 6]. Development of retarded children always needs to be regularly

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monitored via long-term treatment interventions. Whether children receive treatments continuously is important for the success of early intervention.

Treatment for retarded children relies mostly on the parents' ability to meet the needs of treatment. In recent years, a variety of researches have emphasized the importance of parents' participation in their children's intervention processes [7-9]. Professionals in early intervention services are encouraged to understand parents' psychosocial problems and further enhance their capacity regarding care for their retarded children. It seemed worth investigating how the psychosocial factors impact on the continuity of their children receiving early intervention treatment.

The purposes of this study were to identify parents' psychosocial problems that related to care for developmentally delayed children and to find whether the children had continuously received treatments. The second purpose was to find parents' psychosocial variables that could be used to significantly determine whether retarded children would continuously receive treatment programs.

MATERIALS AND METHODS

A questionnaire investigating parents' psychosocial factors was designed for parents to fill out and was implemented by the authors in this study. The draft of the questionnaire was given to ten parents in a preliminary trial. The draft was reviewed for clarity and simplicity based on responses from these parents. The questionnaire was revised to ensure that all meaningful items were included and that the words were easy to understand.

The questionnaire consisted of two main parts.

An introductory paragraph described the purpose of the research and the first part included questions on demographic data of parents and their children. The second part consisted of twenty descriptive items concerning parents' psychosocial conditions. A fourpoint Likert scale (1=not likely, 4=very likely) was used to rate the likelihood for each descriptive item. Another item asked whether the child had received early intervention treatments continuously since diagnosed with developmental delay, and parents needed to respond either yes or no to this item. In this study, early intervention treatments were expressed as rehabilitation programs or special education lessons. Children who had received treatments continuously were represented as having received treatments regularly per week or per month; only occasionally were they sick and absent from treatments.

Subjects were the parents of children who had visited the Joint Evaluation Team for Children with Developmental Delay of Kaoshiung Medical University affiliated hospital in the period from April 1997 to February 1999. Questionnaires were given to the parents on the conditions that: a) parents and their children lived in the region of Kaoshiung city or Kaoshiung county, and b) children had received assessments at the Joint Evaluation Team and were identified as developmental delay. Explanations were given to participant parents for any questions about filling out the questionnaires. The completed questionnaires were collected by mail.

Responses were coded and analyzed using the statistical software package SPSS 8.0 for Windows Applications. Of the responses to items regarding the psychosocial conditions of the parents, factor analysis [10] was used for grouping the items (variables)

Table 1. Demographic data of the participant parents and their developmentally delayed children (N=207)

Items	Mean ± SD	No. of fathers	No. of mothers
Age of fathers (year)	35.9 ± 4.9		
Age of mothers (year)	32.7 ± 4.9		
Age of their children (year)	4.6 ± 2.5		
Educational level of parents:			
Primary school	·	1	7
Junior high school		4	28
Senior high school		11	102
College		7	43
Graduate school		1/ /	3

into a few factors by analyzing the correlation of interrelated quantitative variables. Discriminant analysis [11] was used for investigating and finding out variables that are important for distinguishing between children who had continuously received treatment and those who had not. The internal consistency of these items was measured by examining the strength of correlation among items. Statistical significance was determined based on $\alpha = 0.05$.

RESULTS

A total of 520 questionnaires were given out.

Parents filled out the questionnaires from January 1999 to April 1999. Parents of 227 children (42.5%) returned the questionnaires. Of these returned questionnaires, 207 had been completely finished and these questionnaires were adopted for further statistical analysis. Contact was made with the parents who did not respond. They did not respond because they were too busy, the parents had a negative attitude to questionnaires, of they promised to answer but never did. The demographic data of the 207 participant parents and their delayed children are listed in Table 1.

In the factor analysis for the items of psychoso-

Table 2. Six extracted factors and their factor loadings from the original items regarding parents' psychosocial conditions

Renamed factors and their original items included	Factor loadings	
(Factor 1) Family income		
Poor family economics due to needs of the retarded child	.837	
A great outlay for the child's therapies and education	.818	
Daily life disrupted when caring for the retarded child	.555	
Poor living environment for the retarded child	.483	
Job discontinued due to caring for the retarded child	.425	
(Factor 2) Family members relationship		
Care for the child causing inharmonious spouse relationship	.758	
Family members dispute how to care for the child	.648	
Worry about family members who don't accept the retarded child	.599	
Siblings don't like the retarded child	.428	
(Factor 3) Caring skills		
Feel lack of good caring skills	.688	
Feel lack of adequate caring knowledge	.638	
(Factor 4) Supporting resources		
Don't understand the developmental problems of the child	.686	
Unaware of the services of early intervention	.401	
Doubts about the effects of intervention treatments	.379	
Don't know what social welfare or finance are available	.366	
(Factor 5) Acceptance of the retarded child		
Can't accept the retarded conditions of the child	.593	
Worry about child's delay being known by neighbors or friends	.463	
(Factor 6) Self- psychological adaptation		
Time for rest and self-recreation is decreased	.440	
Care for child causes great psychological stress	.368	
Hard to explain the child's problem to the family	.302	

Table 3. Standardized discriminant function coefficients between variables and discriminant functions (dependent variable)

Variables	discriminant coefficients		
(Factors)	Standardized	Non-standardized	
Family income	0.858 *	0.949*	
Supporting resources	-0.368	-0.452	
Self-psychological adaptation	0.359	0.469	
Caring skills	0.129	0.155	
Family members relationship	-0.076	-0.086	
Acceptance of the retarded child	0.042	0.057	

dependent variable: Children had received treatment continuously (coding 0), Children hadn't received treatment continuously (coding 1)

Table 4. Frequency and percentage analysis for the classification results of discriminant analysis (valid cases=188)

	Predicted groups: subjects number		
	Continuously receiving treatments	Non continuous treatments	
Actual groups			
Continuously receiving treatments	58 (60.4%)	38 (39.6%)	
Non continuous treatments	37 (40.2%)	55 (59.8%)	

cial conditions, after the principal component analysis solution with varimax rotation, six factors were extracted and they accounted for 46.9% of the variance of the twenty items. The six factors were renamed respectively, according to the original items they included. Table 2 lists these six renamed factors, the original items included in each factor, and the factor loading. The reliability coefficient of Cronbach α , for analyzing internal consistency of the items, was calculated as 0.89.

There were 96 positive responses to the item of whether the child continuously received treatment, and there were 92 negative responses (19 missing values), so that about half of the subjects' children had received treatment continuously. This item was also used as a dependent variable for the subsequent discriminant analysis.

The result of discriminant analysis is shown in Table 3. The variables were sorted by values of the

discriminant coefficients. Larger discriminant coefficients of the corresponding variables were thought to contribute more to the overall discriminant function (dependent variable). The first variable listed in Table 3 was found to have statistically significant meaning. Summary classification results of discriminant analysis are listed in Table 4. The correct classification into two predicted groups by this significant variable is 60.4% of the cases. A predicted group was expressed as the most likely group a case would fall into, based on the discriminant analysis, whereas an actual group is simply the group to which a case actually belongs.

DISCUSSION

In the present study, the findings about parents' psychosocial factors extracted in this study were similar to some other researches [12,13]. In a study in-

^{*} p<0.05: family income had a statistically significant contribution to the dependent variable

vestigating 297 parents of intellectually disabled children [12], the results revealed that the parents' needs are information support, professional support, services support, economic support and psychic support respectively. Similarly, in the study of Bailey and Blasco [13], the domains used for assessing family needs are information, support, explaining to others, community services, financial needs and family functioning. Therefore, the psychosocial factors studied in this study were the quite important factors to be regarded when parents' problems are investigated.

The other finding in the present study was that only about half of the patients responded positively that their children had received treatment continuously. From the results of discriminant analysis, family income was found as the significant variable for discriminating whether the children had continuously received treatments or the treatments had been discontinued. We noted that low family income might significantly prevent retarded children from receiving treatments continuously. This result showed that family income was crucial to parents' capacity to meet the needs for receiving treatment. In the present study, beside the factor of family income, other parents' psychosocial factors were not found as significant contributors. We conclude that some other factors that contribute to the continuity of early intervention treatments have still not been identified. In future studies, it is worth investigating other important variables that would influence the continuity of treatments. There is a scarcity of other researches that discuss how family income or financial support would influence developmentally delayed children receiving treatment. This study was to identify significant variables that were important for influencing the continuity of treatment in an attempt to cope with the special problems that parents have. Professional understanding of the problems of parents may greatly assist in providing intervention strategies and services that are meaningful to the parents. Service providers would instruct parents in effective home care skills to reduce transport and time demands on parents visiting treatment departments. Service providers would also supply related knowledge of social financial welfare applications in order to ensure developmentally delayed children could acquire necessary continuous early intervention treatments.

Future researches should be directed at studying effective strategies that can be used to promote necessary continuous treatments in children with developmental delay.

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REFERENCES

- Semmler CJ. Developmental disability. In: Early occupational therapy intervention (Semmler CJ, Hunter JG eds.), Aspen, Maryland, 1988; 232-233.
- Stephens LC, Tauber SK. Early intervention and preschool programs. In: Occupational therapy for children (Pratt PN, Allen AS eds.), CV Mosby Company, New York, 2nd ed., 1989; 382-395.
- 3. Simeonsson RJ, Cooper DH, Scheiner AP. A review and analysis of the effectiveness of early intervention programs. Pediatrics 1982; 69: 635-641.
- Berlin LJ, Brooks-Gunn J, McCarton C, McCormick MC.
 The effectiveness of early intervention: examining risk factors and pathways to enhanced development. Preventive Med 1998; 27: 238-245.
- Wang WY, Chang JJ. A study for problems of parents' involvement in home treatment program for their handicapped children. J Phys Ther Assoc ROC 1996;
 21: 215-221. [In Chinese; English abstract]
- Bailey DB, Simeonsson RJ, Winton PJ, Huntington G, Comfort M, Isbell P, O'Donnell K, Helm J. Family-focused intervention: A functional model for planning, implementing, and evaluating individualized family services in early intervention. J Div Early Child 1988; 10:156-171.
- 7. Chiarello L, Effgen S, Levinson M. Parent-professional partnership in evaluation and development of individualized family service plans. Pediatr Phys Ther 1992; 4:64-69.
- Martin T, Vandeveer M, Rio MD. Family interaction and motor development of children who required intensive care at birth. Pediatr Phys Ther 1992; 4: 78-84.
- Wendt LV, Ekenberg L, Dagis D, Janlert U. A parentcentered approach to physiotherapy for their handicapped children. Dev Med Child Neurol 1984; 26: 445-448.
- Factor analysis. In: SPSS Base 8.0 Application Guide, SPSS Inc., Chicago, 1998; 317-324.
- 11. Discriminant analysis: In: SPSS Base 8.0 Application Guide, SPSS Inc., Chicago, 1998; 243-250.

- 12. Wang TM. Assessing needs of families with young intellectually disabled children. Bulletin Special Educ ROC 1993; 9: 73-90. [In Chinese; English abstract].
- Bailey DB, Blasco PM. Parents' perspectives on a written survey of family needs. J Early Intervention 1990; 14: 196-203.

父母親的心理社會問題對發展遲緩兒童 持續接受早期介入治療的影響

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