

Oifig an Cheannaire Oibríochtaí,

Na Seirbhísí Míchumais/An Rannán Cúram Sóisialta, 31-33 Sráid Chaitríona, Luimneach.

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1st December 2021

Deputy Mark Ward, Dail Eireann, Leinster House, Kildare Street, Dublin 2. e-mail: <u>mark.ward@oireachtas.ie</u>

Dear Deputy Ward,

The Health Service Executive has been requested to reply directly to you in the context of the following parliamentary questions, which were submitted to this department for response.

PQ 56856/21

To ask the Minister for Health the number of children waiting on an audiology service; the average waiting time after the completion of an assessment of needs by CHO area; and if he will make a statement on the matter.

PQ 56857/21

To ask the Minister for Health the average number of interventions received after a child is identified as needing an audiology service after completion of an assessment of needs since their initial intervention in the first 12 months by CHO area in the past five years in tabular form; and if he will make a statement on the matter.

PQ 56881/21

To ask the Minister for Health the number of children waiting on a service for the deaf and hard of hearing; the average waiting time after the completion of an assessment of needs by CHO area; and if he will make a statement on the matter.

PQ 56882/21

To ask the Minister for Health the average number of interventions received after a child is identified as needing a service for the deaf and hard of hearing after completion of an assessment of needs since their initial intervention in the first 12 months by CHO area in the past five years in tabular form; and if he will make a statement on the matter.

PQ 56883/21

To ask the Minister for Health the number of children waiting for appointments in services for the deaf and hard of hearing: the average waiting times by CHO area in tabular form; and if he will make a statement on the matter.

HSE Response

The HSE acknowledges the challenges in meeting the demand for children's disability services and is acutely conscious of how this impacts on children and their families.



Audiology and services are provided mainly through Acute services and Primary Care Teams in CHO Areas to children and young people requiring assessment, diagnosis and treatment. Services for the deaf and hard of hearing are provided in Acute, Primary Care Services and Disability Services

Whilst accepting that significant levels of vital services including audiology and services for the deaf and hard of hearing were temporarily curtailed due to the current Covid-19 pandemic, the restoration and continuity of services is underway in a safe way in line with the very significant investment made by the State and funded agencies. We continue to work with service users and their families/carers to ensure that we achieve this aim.

The information regarding the number of Assessment of Need reports completed since 2017 is provided in Table 1 below. The detail regarding the number of these reports that identified a requirement for audiology and requirement for services for deaf and hard of hearing is also provided in the tables below. This data is not available by CHO area.

HSE Disability Services does not gather data on the service provided through the Acute Hospital sector or Primary Care.

Of note, children do not require an Assessment of Need under the Disability Act to access any health services including audiology and services for deaf and hard of hearing. Children requiring Audiology Services would normally access such services via their GP.

Table 1											
Year	AON Reports Completed	No. with identified need for Audiology	No. with identified need for Services for Deaf and Hard of Hearing								
2017	3660	132	5								
2018	4237	134	4								
2019	3312	89	4								
2020	3911	76	5								
2021	6204	94	1								

Primary Care Service collects data on the number of children waiting on an audiology service and have provided us with this information. Please see Appendix 1- See Table 2 Audiology - Number of 0-17 years & 11 months old Patients waiting for treatment October 2021

Primary Care and Disability Services do not collate information regarding waiting lists for a service for the deaf and hard of hearing.

Progressing Disability Services for Children & Young People (PDS) programme

A number of service improvements are being introduced that, when implemented, will help improve access to services including therapy services for children with disabilities and developmental delays.

The implementation of the Progressing Disability Services for Children & Young People (PDS) programme is agreed Government and HSE policy. This policy supports the reconfiguration of children's disability services across all statutory and non statutory organisations into Children's Disability Network Teams (CDNT) to provide equitable, child and family centred services based on need rather than diagnosis, where the child lives or goes to school. Each Community Healthcare Organisation are working with key stakeholders in their area including parents and education services to ensure that appropriate arrangements are put in place to facilitate in-reach services to special school settings as appropriate to the needs of the children accessing services.

The Progressing Disability Services for Children and Young People (PDS) model addresses the previous inequity in service provision whereby there may have been an excellent service for some children and little or no service for others. This variance may have been linked to diagnosis, age group or geography. Under the PDS programme children's disability services are changing from diagnosis based to needs based, so that all children with a disability or developmental delay have access to the right service based on their needs no matter where they live.



The National Policy on Access to Services for Children & Young People with Disability & Developmental Delay policy provides a single point of entry, signposting parents and referrers to the most appropriate service (Primary Care for non-complex functional difficulties and Children's Disability Network Teams for complex functional difficulties).

The HSE is committed to the full implementation of the Progressing Disability Services for Children and Young People Programme (PDS). PDS is a significant change programme for the provision of services and supports for children from birth to 18 years of age, in line with Sláintecare and the Programme for Government, in order to:

- Provide a clear pathway and fairer access to services for all children with a disability
- Make the best use of available resources for the benefit of all children and their families
- Ensure effective teams are working in partnership with families and with education staff to support children with a disability to reach their full potential.

PDS aligns with two clear objectives of The Sláintecare Report to:

- Provide the majority of care at or as close to home as possible
- Create an integrated system of care with healthcare professionals working closely together.

Children's Disability Network Teams (CDNTs)

In line with the PDS model, resources assigned to children's disability services are allocated to 91 Children's Disability Networks (CDN) aligned to 96 Community Healthcare Networks (CHNs) across the country and each Children's Disability Network will have one CDNT providing services and supports for children aged from birth to 18 years of age. Early Intervention Teams and School Age Teams already in place are reconfiguring into birth-18 CDNTs. On full reconfiguration of children's disability services into CDNTs in December 2021, every child across the country with complex needs arising from their disability will have access to a CDNT

CDNTs are teams of health and social care professionals, including nursing, occupational therapy, psychology, physiotherapy, speech and language therapy, social work and others. The team works closely together in a family centred model, focusing on the child's and family's own priorities. Once all teams are in place, every child with complex needs will have access to a team, regardless of the nature of their disability, where they live, or the school they attend.

Children and their families will have access to the full range of services and supports of the CDNT according to their individual needs and available resources. This includes universal, targeted and specialist supports, such as individual therapeutic intervention and access to specialist consultation and assessment when needed. Supports will be provided as is feasible in the child's natural environments - their home, school and community.

Services in most parts of the country have now reconfigured to CDNTs. The remaining teams in will be established before the end of 2021 thereby, all 91 CDNTs will be in place and delivering services at that time.

Yours sincerely,

Regar

Bernard O'Regan Head of Operations - Disability Services, Community Operations



Appendix Table 2

Aud	Audiology - Number of 0-17 years & 11 months old Patients waiting for treatment													
October 2021														
	0 - 4 Yrs 11 Months				5 - 17 Yrs 11 Months									
CHO Area	0 - ≤ 12 weeks	>12 weeks - ≤ 26 weeks	>26 weeks but ≤ 39 weeks	>39 weeks but ≤ 52	> 52 weeks	0 - ≤ 12 weeks	>12 weeks - ≤ 26 weeks	>26 weeks but ≤ 39 weeks	>39 weeks but ≤ 52	> 52 weeks	TOTAL			
CHO 1 Total	65	56	24	12	6	65	53	17	12	91	401			
CHO 2 Total	101	150	101	98	152	133	154	61	41	64	1,055			
CHO 3 Total	59	79	59	49	139	33	39	29	37	151	674			
CHO 4 Total	239	204	171	173	480	190	156	82	66	74	1,835			
CHO 5 Total	154	172	107	109	252	101	105	75	112	511	1,698			
CHO 6 Total	0	0	0	0	0	0	0	0	0	0	0			
CHO 7 Total	129	128	83	26	0	132	149	68	27	0	742			
CHO 8 Total	102	119	67	47	188	54	31	43	51	605	1,307			
CHO 9 Total	192	143	106	39	0	82	58	68	55	2	745			
Natio nal Total	1,041	1,051	718	553	1,217	790	745	443	401	1,498	8,457			

