

***Original article***

**Title:** Association of maternal age with type of congenital heart disease in patients with Down syndrome: A single-center study.

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## **Abstract**

**Background:** Congenital heart disease (CHD) is commonly seen in patients with Down syndrome (DS) and is closely associated with morbidity and mortality. Maternal age is an established risk factor for DS.

**Objectives:** The aim of the present study is to analyze the incidence of CHD in Down syndrome and the effect of maternal age on this incidence and the type of CHD.

**Materials and Methods:** Six hundred and thirty-one patients with DS who were admitted to pediatric cardiology department, between December 2014 and December 2018, were retrospectively analyzed. The detected CHDs were evaluated in association with maternal age.

**Results:** Four hundred forty-five (70.5%) cases had one or more congenital heart defect. Two hundred and seven patients (32.8%) had isolated simple defect while the remaining had complex or associated multiple CHDs (n:238, 37.7%). Considering all patients, secundum ASD (42.1%) was the most frequent CHD followed by PDA (33.9%), VSD (28.6%) and AVSD (9.6%). Regarding the maternal age, the incidence of CHD was lowest in babies whose maternal age were between 25-35 years. AVSD, was most frequent in patients born to mothers aging  $\leq 20$  years. Incidence of ASD did not markedly differ between maternal age groups while nearly half of the patients born to mothers aging between 21-25 years had PDA and incidence of VSD was markedly increased with the maternal age of  $>45$  years.

**Conclusion:** Distribution of CHD also varies in accordance with maternal age. Babies born to mothers aged  $<25$  or  $>35$  years are more likely to have CHD. Incidence of AVSD, which had been reported to be the most common CHD in patients with DS, has been decreased in time and tends to be mostly associated with maternal age of  $\leq 20$  years.

**Keywords:** *Down syndrome, Congenital heart disease, maternal age.*

## **Introduction**

Down Syndrome (DS) is the most common autosomal chromosome anomaly and closely associated with congenital heart disease (CHD) [1]. CHD is an important feature of these patients which plays an important role on morbidity and mortality [2]. There are studies reporting differences in type of CHD in various geographical areas [3-5]. Maternal age is a well-known risk factor associated with DS [5, 6]. To our knowledge there is not any study in the literature that focuses on the relation between the type of the CHD and the maternal age. The aim of the present study is to analyze the types of CHD and the association of CHDs and maternal age in this special group of patient.

### **Patients and methods**

Six hundred and thirty-one patients with DS who were admitted to pediatric cardiology department at a tertiary center, due to any reason of cardiac evaluation between December 2014 and December 2018, were retrospectively analyzed. Diagnosis of DS were made by either clinical criteria or genetic analysis. Patient demographics, maternal age at delivery and results of echocardiographic records were noted. Patients were divided into 7 groups according to maternal age. Group 1 was consisting of patients with maternal age of equal or less than 20 years while maternal age was 21-25 years, 26-30 years, 31-35 years, 36-40 years, 41-45 years and equal or more than 46 years in group 2, 3, 4, 5, 6 and 7, respectively. The detected CHDs were evaluated in association with maternal age. The congenital heart defects were classified as isolated or associated simple defects (Atrial septal defect (ASD), ventricular septal defect (VSD), patent ductus arteriosus (PDA)), complex defects (Atrioventricular septal defect (AVSD), Tetralogy of Fallot (TOF), double outlet right ventricle (DORV), transposition of great arteries (TGA)) and other defects.

#### *Statistical analysis:*

The statistical analysis was performed using Statistical Package for Social Sciences Software (SPSS), version 21. Demographic and clinical variables were summarized with descriptive statistics. Categorical variables were summarized as absolute frequency and percentage, whereas continuous variables were summarized as median, mean and standard deviation.

## Results

The mean and median age of patients at diagnosis was  $3.32 \pm 4.25$  years and 1.09 years, respectively (range: 0 days-17.41 years). The male-to-female ratio was 1.5/1. Three hundred and sixty patients (57.1%) were under 2 years-old while 147 patients (23.3%) were between 2-7 years and the remaining (n:124, 19.7%) above 7 years. The mean maternal age at delivery was  $34.92 \pm 7.40$  years (range: 17-50 years). Almost half of the mothers were between 31 to 40 years old. The distribution of maternal age is shown on figure 1.

Four hundred forty-five (70.5%) patients had one or more congenital heart defect. Two hundred and seven patients (32.8%) had isolated simple defect while the remaining had complex or associated multiple CHDs (n:238, 37.7%). There was not a significant difference by means of CHD in accordance with gender. Distribution of CHDs detected in study population is shown on table 1. Considering all patients, secundum ASD was the most frequent CHD, but the vast majority of the patients with secundum ASD were under 2 years and most of secundum ASDs were small defects which were expected to have spontaneous closure with somatic grow-up (Figure 2). ASDs were isolated single defects in 91 patients (14.4%) whereas, the remainders (n:175, 27.7%) were in association of various defects. Sinus venosus type ASD was not detected in any of patients, while primum type ASD was detected only as a component of AVSD. PDA was the second most common CHD (33.9%). PDAs in patients under 3-month old were not included. Hemodynamically important PDAs were consisting 74.3% of all PDAs so a transcatheter closure was planned or performed while the remaining had silent or small ducts without hemodynamic importance. Isolated PDA as a single defect was accounting 9.8% of all subjects. VSD was the third most common defect (28.6%). Isolated single VSD was found in 8.7% of all study population and perimembranous VSD was the most common type. AVSDs were detected in 61 cases (9.6%). The most common form was Rastelli type C, and the majority had balanced ventricles. The majority of the AVSDs were in association with other CHDs, whereas isolated AVSD was accounting for 2.2% of all cases. More than half of the patients with CHD had two or more associated defects (53.4%). Association of ASD and PDA was the

most frequent association of CHD followed by VSD with PDA and ASD (Table 2). Tetralogy of Fallot was detected in 12 patients (1.9%) and was the most common cyanotic CHD.

Regarding the maternal age, incidence of CHD was lowest in babies of whom maternal age were between 25-35 years. Mothers who were below 25 years or above 35 years had similar incidence of having a child with CHD (Figure 3). There were differences in types of CHDs between groups (Figure 4). ASD was the most common CHD in all groups except in group 2, in which PDA was the most common CHD. VSD was the most common CHD in group 7 with almost the same percentage with ASD in this group. AVSD, which has been reported to be the most common defect in various studies in patients with DS, was most frequent in group 1 (33.3%). Its incidence markedly decreased in patients born to mothers between 21-to-30 year-old and reached an account of %11.4-11.9 in babies born mothers older than 40 years (Figure 5).

## Discussion

Congenital heart disease is a common cause of mortality and morbidity in patients with DS and it is reported in various studies that 44 to 79.2% of these patients have CHD [7-10]. The overall incidence of CHD within our study population was 70.5%. This rate was similar to that studies reported from different regions of our country [10, 11] but higher than some recently reported studies in the literature [12, 13].

The reason for the high incidence seen in our study could be that minor lesions such as small secundum ASDs in infants and young children as well as small PDAs without audible murmur which do not have hemodynamic importance were included. The incidence would be lower if these lesions were excluded.

Although varying studies have reported that the distribution of CHDs in DS may vary according to ethnicity, in the majority of the studies AVSD, VSD, and ASD are reported to be still the most common three types of CHD in this group of patients [6, 9, 14]. In western European countries and the USA, AVSD (43%) is reported to be the most common CHD associated with DS followed by, VSD (32%); secundum atrial septal defect (10%); tetralogy of Fallot (6%); and isolated PDA (4%) and about 30% of patients have several cardiac defects [7, 15]. However, in Asia, isolated VSDs have been reported to be the most common defect, observed in about 40% of patients [16], whereas in most reports from Latin

America, the secundum type of ASD is suggested to be the most common lesion [4, 17, 18]. Kim et al.[12] reported ASD as the most common CHD, occurring in 30.5% of all DS, followed by VSD (19.3%), PDA (17.5%), and AVSD (9.4%) in Korean patients. In two studies from different regions of Turkey, Gul et al. [10] have reported ASD as **the** most common CHD in patients with DS whereas, Nisli et al. [11] found AVSD as the leading CHD in this group. Vida et **al.** reported a high frequency of PDA [18]. In that study, PDA was the most common single defect and was also the most frequent concomitant malformation. Pfitzer et al. [14] have examined the dynamics in the prevalence of patients with CHD and DS over an extended period and reported that the prevalence of AVSD and complex CHD have decreased while simple defects such as ASD and VSD are increasing over the time. Similar to that, Bergström et al. [5] have reported temporal changes with regard to specific defects during a 20-year period. They found that AVSD and VSD were accounting for 46% and 14% of CHD, respectively, in the first periods of the study while this amount was shifted to 31% and 30%, respectively, in the last periods of the study. They suggested that the overall incidence of congenital heart defects among newborns with DS remained stable over the time while the risk of complex malformations declined and simple defects like VSD and ASD increased. In our study, ASD was the most common CHD, while AVSD was lesser when compared with the majority of previously reported studies, but closely similar to the Korean study [12].

High maternal age is a well-known risk factor for meiotic non-disjunction and still the only established risk factor in the etiology of DS [19, 20]. Regarding the prevalence of CHD in infants with DS in accordance with maternal age, Kim et al. [12] and Scott et al. [9] have reported that mothers  $\geq 35$  years are more likely to give birth to a DS child with a CHD. However, there are studies which reports the increased incidence of DS in young mothers [21, 22]. Chaohab et al. reported **that** maternal age of under 32 years has a higher risk for CHD **in patients with DS** [23]. **Bergström et al. [5]. reported that the incidence of any CHD in DS is most common in patients with maternal age of under 25 years. The incidence is reported as 62.5% if maternal age is below or equal to 19 and 58.0% if maternal age is between 20-25 years.** In the same study the incidence of any CHD was found to be between 53.9-56.4 in babies born to mothers aging 25-35 years, and 50-51% in babies born to mothers aging above 35 years. In our study, maternal age of

between 25 and 35 years was found to be less associated with having CHD while maternal ages of <25 years or >35 years had a higher incidence of having CHD in patients with DS. Incidence of ASD did not markedly differ between maternal age groups while nearly half of the patients born to mothers aging between 21-25 years had PDA and incidence of VSD was markedly increased with the maternal age of >45 years. Interestingly, AVSD was mostly found in babies of young mothers aging 20 years or below with an incidence of more than three times to other maternal age groups.

## **Conclusion**

This retrospective study confirms that the type of CHDs in DS varies with geographical areas and ethnicity. Distribution of CHD also varies in accordance with maternal age. Babies born to mothers aged <25 or >35 years are more likely to have CHD. Incidence of AVSD, which had been reported to be the most common CHD in patients with DS, has been decreased in time and tends to be mostly associated with maternal age of  $\leq 20$  years.

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**Authors' Contributions:** 'Epçaçan S' designed the study, managed the literature searches, wrote the protocol, and wrote the manuscript. 'Tunçdemir P and 'Karakuş-Epçaçan Z' managed the analyses of the study and language editing of the manuscript. 'Tunçdemir Y' performed the statistical analysis. All authors read and approved the final manuscript.

**Ethical approval:** All procedures performed were in accordance with the ethical standards and with the 1964 Helsinki Declaration and its later amendments. Institutional ethical committee approved the study.

## **References:**

1. Weijerman ME, de Winter JP. Clinical practice. The care of children with Down syndrome. *Eur J Pediatr*. 2010 Dec;169(12):1445-52. PMID: 20632187.
2. Vis JC, Duffels MG, Winter MM, Weijerman ME, Cobben JM, Huisman SA, et al. Down syndrome: a cardiovascular perspective. *J Intellect Disabil Res*. 2009 May;53(5):419-25. PMID: 19228275.
3. Narayanan DL, Yesodharan D, Kappanayil M, Kuthirolly S, Thampi MV, Hamza Z, et al. Cardiac spectrum, cytogenetic analysis and thyroid profile of 418 children with Down syndrome from South India: a cross-sectional study. *Indian J Pediatr*. 2014 Jun;81(6):547-51. PMID: 23934063.
4. de Rubens Figueroa J, del Pozzo Magana B, Pablos Hach JL, Calderon Jimenez C, Castrejon Urbina R. Malformaciones cardiacas en los ninos con sindrome de Down. *Rev Esp Cardiol*. 2003 Sep;56(9):894-9. Spanish. PMID: 14519277.
5. Bergstrom S, Carr H, Petersson G, Stephansson O, Bonamy AK, Dahlstrom A, et al. Trends in Congenital Heart Defects in Infants With Down Syndrome. *Pediatrics*. 2016 Jul;138(1): pii: e20160123. PMID: 27252035.
6. Benhaourech S, Drighil A, Hammiri AE. Congenital heart disease and Down syndrome: various aspects of a confirmed association. *Cardiovasc J Afr*. 2016 Sep/Oct;27(5):287-90. PMID: 27805241.
7. Freeman SB, Taft LF, Dooley KJ, Allran K, Sherman SL, Hassold TJ, et al. Population-based study of congenital heart defects in Down syndrome. *Am J Med Genet*. 1998 Nov 16;80(3):213-7. PMID: 9843040.
8. Paladini D, Tartaglione A, Agangi A, Teodoro A, Forleo F, Borghese A, et al. The association between congenital heart disease and Down syndrome in prenatal life. *Ultrasound Obstet Gynecol*. 2000 Feb;15(2):104-8. PMID: 10775990.
9. Scott C, Thame M. The Incidence of Cardiac Lesions among Children with Down's Syndrome in Jamaica - A Prospective Study. *West Indian Med J*. 2014 Dec;63(7):693-7. PMID: 25867555.
10. Nisli K, Oner N, Candan S, Kayserili H, Tansel T, Tireli E, et al. Congenital heart disease in children with Down's syndrome: Turkish experience of 13 years. *Acta Cardiol*. 2008 Oct;63(5):585-9. PMID: 19014001.
11. Gul O GD, Dogan M. Down sendromlu çocukların klinik ve ekokardiyografik değerlendirmesi. [Article in Turkish]. *Turkish J Pediatr Dis*. 2016 (2):116-9. Turkish. DOI: 10.12956/tjpd.2016.263.
12. Kim MA, Lee YS, Yee NH, Choi JS, Choi JY, Seo K. Prevalence of congenital heart defects associated with Down syndrome in Korea. *J Korean Med Sci*. 2014 Nov;29(11):1544-9. PMID: 25408587.
13. Bermudez BE, Medeiros SL, Bermudez MB, Novadzki IM, Magdalena NI. Down syndrome: Prevalence and distribution of congenital heart disease in Brazil. *Sao Paulo Med J*. 2015 Nov-Dec;133(6):521-4. PMID: 26648279.
14. Pfitzer C, Helm PC, Rosenthal LM, Berger F, Bauer UMM, Schmitt KR. Dynamics in prevalence of Down syndrome in children with congenital heart disease. *Eur J Pediatr*. 2018 Jan;177(1):107-15. PMID: 29127498.
15. Laursen HB. Congenital heart disease in Down's syndrome. *Br Heart J*. 1976 Jan;38(1):32-8. PMID: 1252293.
16. Jacobs EG, Leung MP, Karlberg J. Distribution of symptomatic congenital heart disease in Hong Kong. *Pediatr Cardiol*. 2000 Mar-Apr;21(2):148-57. PMID: 10754087.
17. Frid C, Drott P, Lundell B, Rasmussen F, Anneren G. Mortality in Down's syndrome in relation to congenital malformations. *J Intellect Disabil Res*. 1999 Jun;43 ( Pt 3):234-41. PMID: 10392609.
18. Vida VL, Barnoya J, Larrazabal LA, Gaitan G, de Maria Garcia F, Castaneda AR. Congenital cardiac disease in children with Down's syndrome in Guatemala. *Cardiol Young*. 2005 Jun;15(3):286-90. PMID: 15865831.
19. Hartway S. A parent's guide to the genetics of Down syndrome. *Adv Neonatal Care*. 2009 Feb;9(1):27-30. PMID: 19212162.
20. Hecht CA, Hook EB. Rates of Down syndrome at livebirth by one-year maternal age intervals in studies with apparent close to complete ascertainment in populations of European origin: a proposed



revised rate schedule for use in genetic and prenatal screening. Am J Med Genet. 1996 Apr 24;62(4):376-85. PMID: 8723068.

21. Adeyokunnu AA. The incidence of Down's syndrome in Nigeria. J Med Genet. 1982 Aug;19(4):277-9. PMID: 6214633.

22. Hoe TS, Boo NY, Clyde MM. Incidence of Down's syndrome in a large Malaysian maternity hospital over an 18 month period. Singapore Med J. 1989 Jun;30(3):246-8. PMID: 2531468.

23. Chehab G, Chokor I, Fakhouri H, Hage G, Saliba Z, El-Rassi I. Cardiopathie congenitale, age maternel et consanguinite parentale chez les enfants avec syndrome de Down. J Med Liban. 2007 Jul-Sep;55(3):133-7. French. PMID: 17966733.

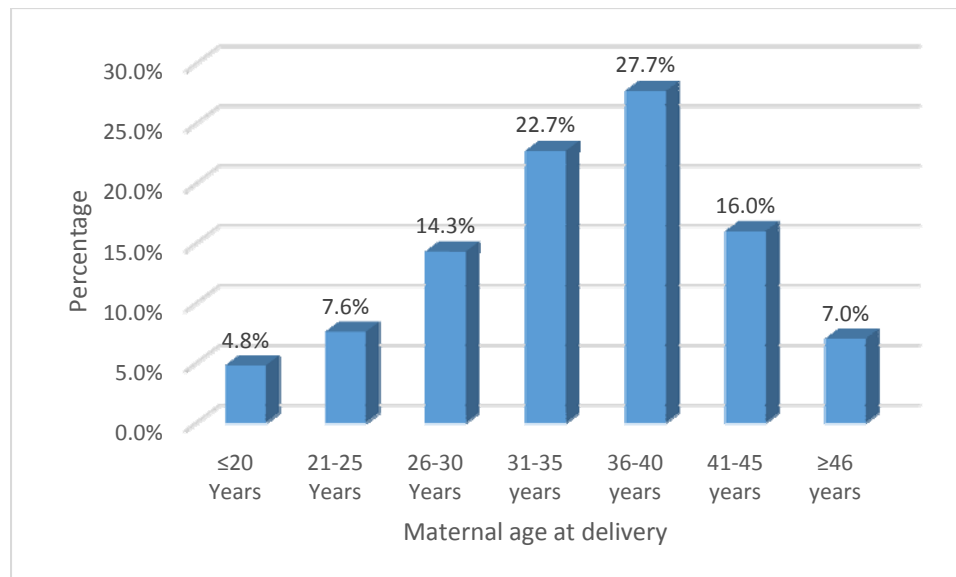


Figure 1: Incidence of any CHD in accordance with maternal age.

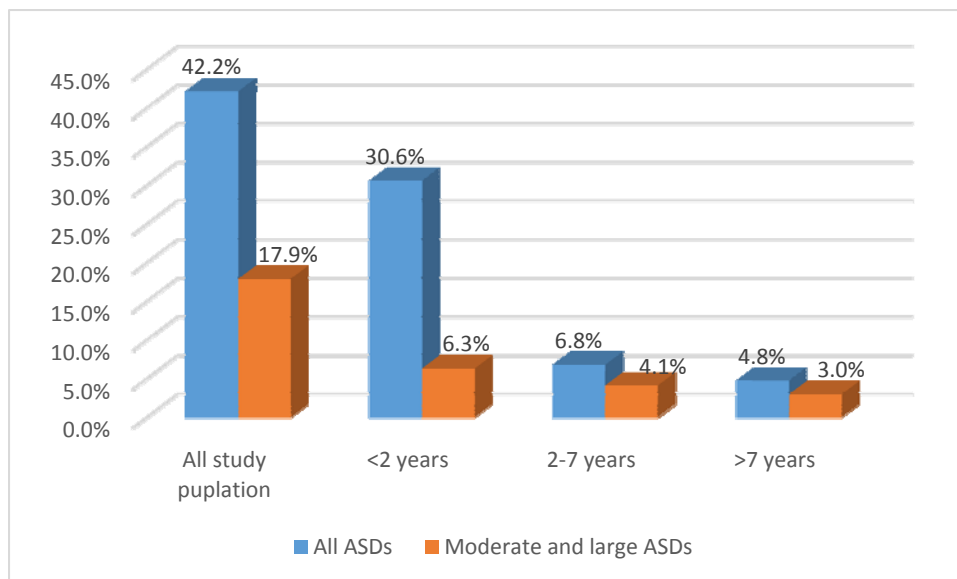


Figure 2: Distribution of ASD by means of defect size in patient age groups

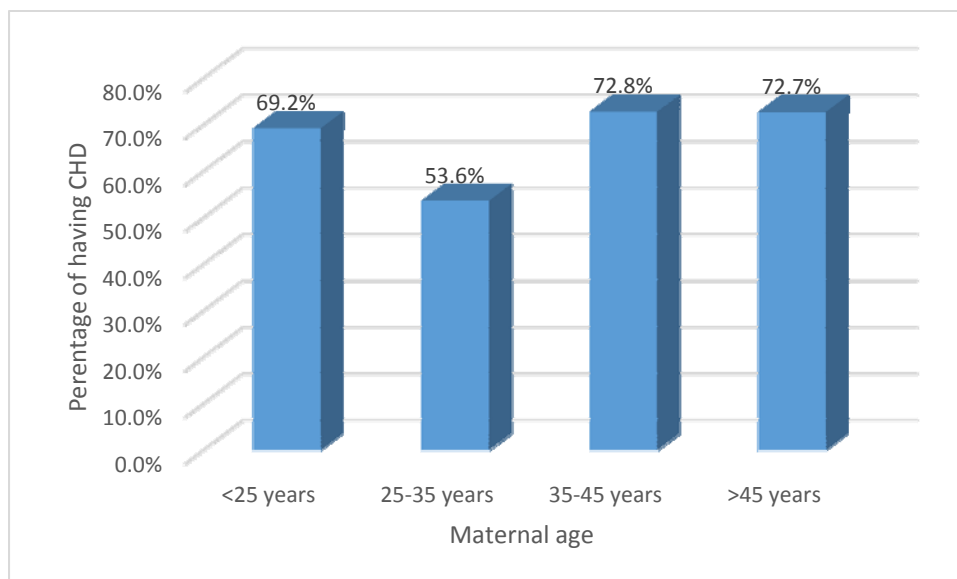


Figure 3: Incidence of CHDs according to maternal age.

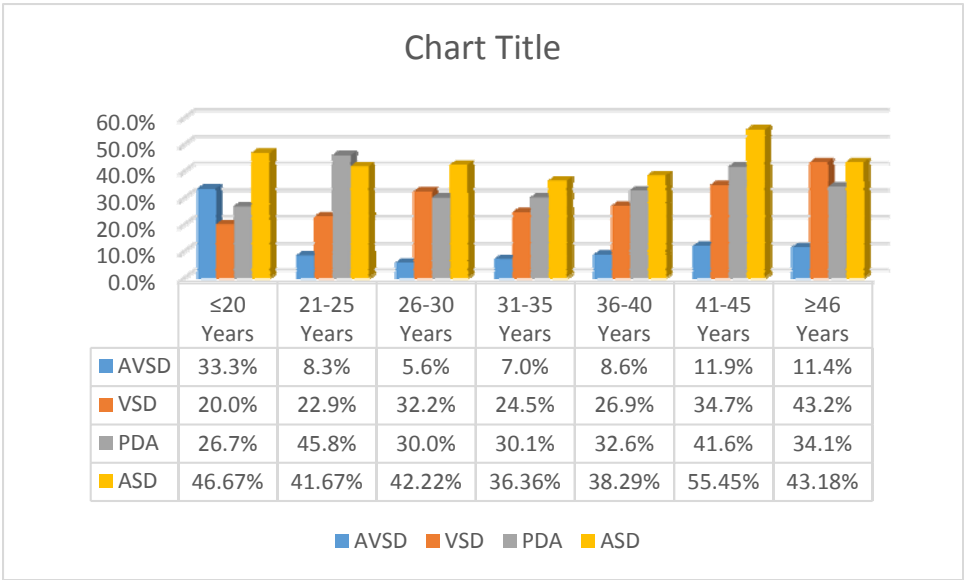


Figure 4: Distribution of mostly seen CHDs in maternal age groups.

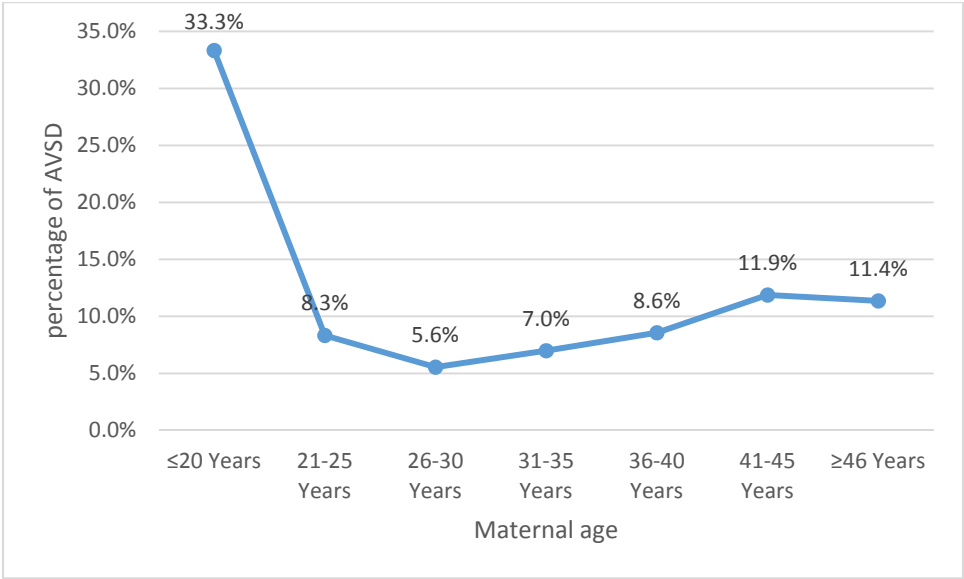


Figure 5: The changing incidence of AVSD in accordance with maternal age.

<b>Table 1:</b> Distribution of CHDs in all patients with DS (n=631)		
<b><i>Congenital Heart Defect</i></b>	<b><i>n</i></b>	<b><i>(%)</i></b>
<b>Atrial septal defect</b>	266	(42.1)
Ostium secundum	266/266	(42.1)
Sinus venosus	0	0
<b>Patent Ductus Aretriosus</b>	214	(33.9)
<b>Ventricular septal defect</b>	181	(28.6)
Perimembranous	131/181	(72.3)
Muscular	34/181	(18.8)
Inlet	16/181	(8.9)
<b>AtRIOventricular septal defect (AVSD)</b>	61	(9.6)
Complete AVSD	57/61	(93.4)
<i>Rastelli type A</i>	15/57	(26.3)
<i>Rastelli type B</i>	2/57	(3.5)
<i>Rastelli type A</i>	40/57	(70.2)
Balanced	53/57	(92.9)
Unbalanced	4/57	(7.1)
Intermediate AVSD	4/61	(6.6)
Partial AVSD	0	0
<b>Tatralogy of Fallot</b>	12	(1.9)
<b>Malpositon of the great arteries</b>	1	(0.16)
<b>Double outlet right ventricle</b>	3	(0.48)
<b>Other</b>		
Left superior vena cava	4	(0.64)
Partial anomalous pulmonary venous connection	2	(0.32)
Coartaction of the aorta	1	(0.16)
Pulmonary stenosis	5	(0.8)
Mitral valvular insufficiency	24	(3.8)
Aorticvalvular insufficiency	22	(3.5)
Mitral valve prolapsus	8	(1.27)
Bicuspid aortic valve	8	(0.95)
Subaortic discrete membrane	1	(0.16)
Small-sized asymptomatic coronary artery fistula	2	(0.32)
Hypertrophic cardiomyopathy	1	(0.16)
Right sided aortic arch	3	(0.48)

<b>Table 2: Types of associated CHD in DS</b>		
<b>CHD associations</b>	<b>n,238</b>	<b>(53.4%)</b>
<b>ASD, PDA</b>	62	13,9%
<b>VSD,ASD,PDA</b>	35	7,9%
<b>VSD,ASD</b>	33	7,4%
<b>VSD,PDA</b>	23	5,2%
<b>AVSD</b>	14	3,1%
<b>AVSD,VSD,ASD,PDA</b>	12	2,7%
<b>AVSD,VSD,ASD</b>	12	2,7%
<b>Tatralogy of Fallot</b>	10	2,2%
<b>AVSD,PDA</b>	9	2,0%
<b>AVSD,ASD,PDA</b>	8	1,8%
<b>VSD,ASD,PS</b>	3	0,7%
<b>Tatralogy of Fallot, ASD</b>	3	0,7%
<b>AVSD, ASD</b>	2	0,4%
<b>ASD, PAPVC</b>	2	0,4%
<b>VSD,PS</b>	2	0,4%
<b>AVSD,VSD,ASD,PS</b>	2	0,4%
<b>AVSD,PS</b>	1	0,2%
<b>AVSD,ASD,PS</b>	1	0,2%
<b>COA, PDA</b>	1	0,2%
<b>AVSD,DORV,d-TGA, PDA,PS</b>	1	0,2%
<b>DORV,VSD,ASD,PS</b>	1	0,2%
<b>AVSD,DORV,PDA,PS</b>	1	0,2%