INTRODUCTION

Down syndrome common chromosomal disorder, related with multiple congenital anomalies with mental retardation in the newborns. Incidence of Down syndrome varies from 1 in 600 to 1 in 1000 in live born infants [1, 2]. In India, incidence of Down syndrome is 1 in 1250 [1].

By the Cytogenetic Analysis the presence of the extra 21 in DS has been classified free Trisomy 21, translocation trisomy 21 and mosaic trisomy 21 [3]. Free trisomy 21 is the most common variety, seen in 95% cases and occurs due to paternal meiotic non disjunction [1], 2 to 4% showed a Robertsonian translocation and 1 to 3% showed mosaicism.

The clinical features are important for an early diagnosis to reduce morbidity and mortality. Apart from karyotype most characteristic features are mental retardation, congenital heart defects, facial features and in some cases developing AML at later stage [4].

The oral motor approached activities are effective in enhancing the feeding performance among children with Down’s syndrome. It was also found that the therapy protocol has a greater effect in reducing tongue thrust thereby improving oral functional skills [5].

The extra chromosome 21 is translocated to the acrocentric chromosome of D group (Chromosome 13, 14, 15) or G group (Chromosome 21, 22). Such translocations are usually Robertsonian in type [1]. Non homologous Robertsonian translocation between chromosome 14 and 21 [rob (14q; 21q)] is the most common type while homologous Robertsonian translocation between chromosome 21 and 21 [rob(21q; 21q)] is the second most common type [6]. Translocation Down Syndrome can be inherited from carrier parents [7].

Down syndrome occurs in all races & economic levels. It is caused by third copy of chromosome 21, there are there forms of DS. Simple Trisomy 21, Translocation Trisomy and Mosaic Trisomy. The aim of the study is to know cause of Down syndrome. Chromosomal analysis was carried out by G banding technique. Materials and Methods: 1 ml of peripheral blood samples were collected in Out Patient Department of pediatrics and Cytogenetic analysis was performed. Results: Out of 28, 3 female cases, 2 male cases were Down syndrome. All the 5 cases were free trisomy 21, which is common type of Down syndrome; we have not identified Robertsonian translocation and mosaic type of DS. Conclusion: The present analysis shows that genetic risk factors are responsible for the incidence of Down syndrome.

Keywords: Down syndrome; North Karnataka; Prevalence; G-banded Karyotype.
rence of different genetic disorders including population growth, demographic pattern, occupational shifts with high proportion of individuals engaged in sedentary lifestyle and substantially higher proportion of pre-obese and obese individuals. Consanguineous marriage is a traditional practice in many communities around the world. By keeping these points, in this study an attempt was made to trace out the role of in Down syndrome in north Karnataka population through Karyotyping Analysis.

MATERIALS AND METHODS

Study design: An observational descriptive study

Ethics approval: Study was approved by the institutional ethics committee and informed consent was taken from the parents.

Study location: Genetics laboratory, department of Anatomy, BLDE (DU), Karnataka

Study period: The study was carried from June 2016 to January 2017.

Inclusion criteria: All the participants were taken from our hospital, children’s of age 0-5 years. Down Syndrome cases were recruited in our study.

Exclusion criteria: Other congenital defects and CHD cases as well as patients from other hospitals were excluded from the study.

Sample size: The present study subjected total of Twenty eight pediatric patients, with the age group of 3 to 120 days old out of which 12 were female and 16 were male.[4]

Methodology:

For Cytogenetic analysis we collected 1 ml peripheral blood samples from Out Patient Pediatric department. Karyotyping was carried out for peripheral lymphocytes, cultured from peripheral blood and stained with Giemsa stain as per the Standard Operating Protocol. Olympus Trinocular Research Florescent In Situ Hybridization (FISH) with Applied Spectral Imaging Karyotyping System (Manufacturer name: Olympus, Japan; model: CH20i) was also used during the study. 20-40 spreads were analyzed for each case. The slides were analyzed for detection of chromosomal abnormality for Down syndrome (Trisomy 21).

RESULTS

A total of 28 clinically suspected cases were referred to our genetics laboratory for cytogenetic analysis. Of which 5 cases were found confirmed DS. The rest 23 cases were found to be other chromosomal anomalies. The maternal age of mother of DS patients are not advanced as shown in (Table 1).The age of all DS patients were calculated and it was found to be 3 days to 4 months as shown in (Table 2). Descriptive Statistics shows age mother with age DS patients in terms of standard deviation mother age will be 0.64 & patients will be 24.6 (Table 3). Proportion rate of Down syndrome is 18%, other syndrome will be 82% (Table 4). All these cases were found to be free trisomy form (Figure 1).

Table 1. Distribution of patients according to Age of mothers

| Age (Years) | Frequency | Percent |
|------------|-----------|---------|
| 24         | 1         | 3.6     |
| 25         | 13        | 46.4    |
| 26         | 13        | 46.4    |
| 27         | 1         | 3.6     |
| Total      | 28        | 100.0   |

Table 2. Distribution of patients according to Age of Infants

| Days | Frequency | Percent |
|------|-----------|---------|
| 7    | 10        | 36      |
| 8-14 | 6         | 21      |
| 14-21| 2         | 7       |
| 22-30| 4         | 14      |
| 30   | 6         | 21      |
| Total| 28        | 100.0   |

Table 3. Descriptive Statistics.

| Age                  | N   | Minimum | Max  | Mean  |
|----------------------|-----|---------|------|-------|
| Age of Mothers       | 28  | 24      | 27   | 25.5±0.64 |
| Age of infants       | 28  | 3       | 120  | 20.7±24.6 |

Table 4. Proportion of Down syndrome

| Diagnosis       | N  | Percentage |
|-----------------|----|------------|
| Down Syndrome   | 5  | 18         |
| Other Syndrome  | 23 | 82         |
| Total           | 28 | 100        |

Figure 1. Down syndrome with free 21 normal trisomy female (47XX)
DISCUSSION

Down syndrome has a prevalence of one in 500 to one in 1,000 live births, due to disorder of development arising from incomplete embryogenesis as a result of an additional chromosome 21 copy in the karyotype. This extra chromosome is derived from an over-expression of genetic material due to a tripling of the number of genes. This phenomenon produces structural and functional disorders of the all the systems of the body [14]. There are three forms of DS namely simple trisomy, Mosaic trisomy and translocation trisomy. Karyotype analysis of our study finds only the simple trisomy, in which a form of faulty division of reproductive cell (generally occurring during 1st or 2nd meiotic division) prior to or at conception that results in an embryo with three copies of chromosome 21. As the embryo develops, the extra chromosome is replicated in every cell of the body. The parental and the meiotic/mitotic origin of the additional chromosome 21 could be determined in the early 1990’s with the help of DNA markers [15]. Simple Trisomy 21 in fact constitutes the most common form of DS in children and occurs at rates of 90-95%. Karyotype for female are 47, XX, +21 and 47, XY, +21 for male [15].

Translocation involves transferring a chromosome fragment to centromere of another acrocentric chromosome. In DS such translocation occurs between chromosomes 14 and 21 which occur at rates of 5-6% of all DS cases. The karyotype for female are 46, XX, der (21;21), +21 or 46, XX, der (14;21), +21 and for male 46, XY, der (21;21), +21 or 46, XY, der (14;21), +21[15].

A seldom form of DS is a condition known as mosaicism, which accounts for only 1-2% of all cases of full trisomy 21 DS Mosaicism characterized by the non-disjunction of chromosome 21 takes place in one of the initial cell divisions after fertilization, resulting in a mixture of two types of cells in the body, some with 46 and some with 47 chromosomes. The physical features of Down’s may be milder in individuals with mosaicism. Trisomy 21, especially if the proportion of normal cells is large [16]. All diagnosed cases were reported in our study were having confirmed DS in north Karnataka region. The frequency of free trisomy reported in the present study is most common seen which is in accordance with earlier reports [17, 18, 19]. Other studies revealed that advanced maternal age combined with advanced maternal age significantly influences the incidences of DS [20]. However, earlier studies in the world reports regarding the maternal and paternal ages that increases the risk of chromosomal aneuploidy [21]. Present study shows that the overall female ratio was more compared to male. Other studies reported that excess of males appears to be universal and reported in literature from all over the world. Present study Translocation trisomy21 & mosaic 21were not observed. Translocation and mosaic which are unique in nature as compared to previous studies [18].

There were reports of older maternal age having DS in previous studies from different countries [22, 23]. On the other hand, few studies have also reported incidences of DS to a much young maternal age [17, 24]. In our study revealed that incidences of DS is young maternal age.

Several studies have been done on incidence of DS, but a complete understanding of the cause is yet to be ascertained. Risk factors of DS unlike maternal age, recently Coppode et al. [9] have suggested folate polymorphism as genetic risk factor for birth of DS child in Caucasian women and Shalaby [11] reported consanguinity, drug and environmental toxins as other risk factors.

CONCLUSION

In the Present study Genetic factors & maternal age are higher incidences in Down syndrome. In the present study conclude that free trisomy 21 is most common type Down syndrome chromosomal abnormality in this region.

Suggestions: It is important to educate & create awareness to family at high risk occurrence to go for screening during pregnancy. It is necessary to adopt appropriate prevention strategies and interventions in high risk individuals to curb the growing epidemic of Down syndrome. Innovative community outreach programmes need to be designed and implemented for creating awareness, early screening and treatment of Down syndrome.

Limitation of study: India is known for high degree of inbreeding with its heterogynous population. This makes it necessary to screen a large number of patients perhaps within each group in order to get a true picture of Down syndrome. In order to find out the prevalence of any risk factors, a large number of families need to be investigated

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Conflict of Interest: Declared none

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