Importance of Medical Genetics Research in Medicine

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Short Communication

About millions of families are affected by hereditary diseases around the world. Statistics analysis showed that about 5% of all pregnancies give a child having different anomalies including a genetic disorder, congenital malformation or significant disability. It is estimated that 43% of patients having mental retardation are due to monogenic disorder or chromosomal anomalies [1].

In developed countries, fully or partially inherited diseases account for approximately 45% of admissions pediatrics in hospitals [1]. In developing countries, perinatal and infant mortality accounts the most important percentage of hereditary diseases reaching 25% [2-4]. The most non-infectious diseases causing death have probably a genetic component in developed countries [5].

Although many of the affected individuals lead normal lives and do not necessarily have observed pain, many individuals and families still touched by genetic diseases despite the quality of their life and the medical supervision by which are surrounded. For example, people with mental retardation and congenital disabilities have almost normal lives in their developed countries. It is their families’ responsibility to take care of them. Statistics analysis realized in the USA estimated that about 2 million people having mental retardation, of whom only 4% live in institutions. Most others live at home.

Many and different medical and social services provided to them represent a significant and an expensive cost, as well as economic loss due to the care for people with genetic diseases provided particularly by their families and the society. In this context, it seems very important to continue efforts to develop medical research with particular attention to medical genetics which is able to provide effective treatments for all health communities.

Medical genetics is the specialty most targeted for providing services to people with genetic diseases and their families [6]. These services ameliorate the life of people having genetic disorders and their families and help them to have breasts children using medical services (diagnosis, treatment, rehabilitation or prevention) and social assistance systems to adapt them to their situation and put them on the page of new developments affecting them [7-11].

The conditions studied by geneticists include monogenic anomalies diseases affected by one defective gene (e.g. hemophilia and cystic fibrosis), diseases caused by the interaction of many genes in relation with environmental factors (e.g. diabetes, hypertension and cardiovascular diseases) and conditions due to aberrations chromosomal (e.g. Down syndrome).

In medical genetics, the work diagnosis includes molecular analyzes (DNA, proteins and chromosomes) and clinical manifestations of conditions, including malformations birth. If monogenic diseases are rare, those caused by the gene/environment interaction are common and include many diseases. Prevention in medical genetics involves for common pathologies the identification of subjects with high risk, with the intention of preventing the disease (e.g. heart disease) or being able to diagnose and treat early (multiple cancers) [12]. At present, efforts to important research aim to develop somatic gene therapies to treat or prevent disease by blockage of their functions [13,14].

Medical genetics services must be established at all levels of health care, under the direction of specialist's primary-care physicians [15]. Different health personnel can intervene depending on the severity of the case and the provision of health care system. The genetic team can be made up of different members including essentially genetic doctors, health professionals, agents trained health counselors or genetic counselors, laboratory staff and of course social workers [14,16].

References