**Ethical Considerations Approval**

The researcher should consider the ethical, legal and regulatory norms and standards for research involving human subjects in their own countries as well as applicable international norms and standards.

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1. The participation in the study was completely voluntary.
2. All the needed information for patients or their parents should be presented by using local and simplified terms for a disease in their common language and invite them to be part of this research. We kept patient information confidential and did not publish patient identity.
3. Any patient can talk to anyone he/she feels comfortable with about the research, ask researcher or physician or medical staff any question about research work sample, the way to gain sample and the purpose of using the sample in the present time or future
4. All clinical trials were conducted according to the [Helsinki Ethical Principles](https://www.wma.net/policies-post/wma-declaration-of-helsinki-ethical-principles-for-medical-research-involving-human-subjects/#:~:text=The%20World%20Medical%20Association%20(WMA,identifiable%20human%20material%20and%20data))
5. Authors have disclosed all their conflict of interests if exist.

Corresponding author can sign in behalf of the participating authors

**Name**

**Affiliation**

**The name of the ethical approval body**

**Signature**

**Date**