

European Standards of Care for Children with Cancer

Report on non-medical aspects of care over
children with cancer



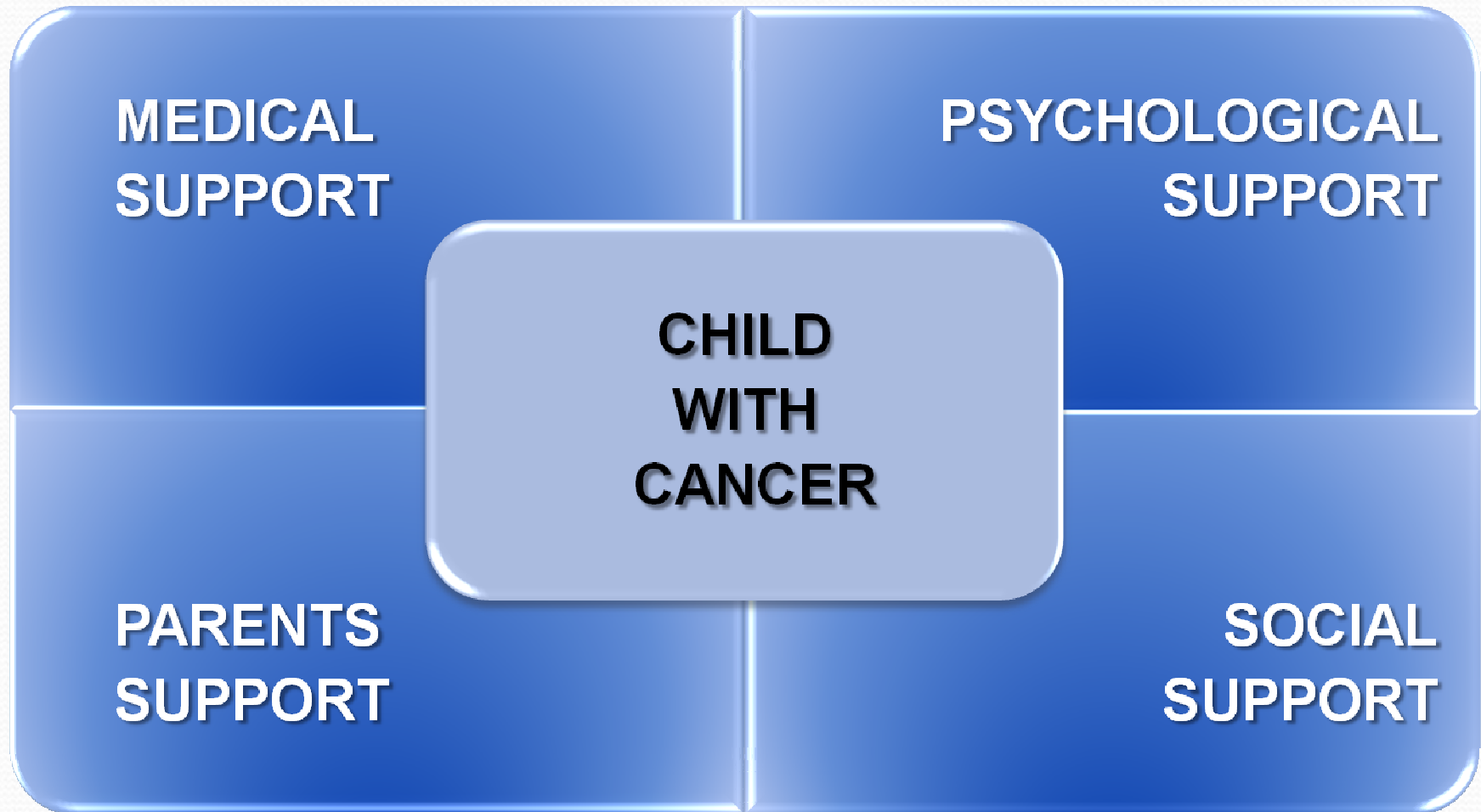
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The Aim

The aim of the report is to present how important role in the process of treatment play non-medical aspects of care over children with cancer.

System of care



The rights of the hospitalized child

- The child has the right to constant parental assistance in the whole process of treatment
- Special conditions should be created for family in order to enable the family to be with the child all the time

Social care

- parents should have an access to information, where they can receive support
- accommodation should be provided for parents
- parents should have an access to support provided by non-profit organisations, voluntary organisations, parents' organisations, non-profit child cancer foundations

The rights of the hospitalized child

- Patients and their parents should receive UNDERSTANDABLE information on the process of treatment
- The child should have the right to play and to education adjusted to his or her age
- The staff should take care of child's development
- The privacy of the child should be respected throughout the process of treatment
- Cultural, language and racial differences should always be respected

Thank you for attention!

Ewa Jack-Górska

