Diploma project, the issue of informal care for a close person with Parkinson's disease from the perspective of caregivers is focused on the intensity of care and access to services that could help informal caregivers in the care of their loved ones. In the theoretical section of Parkinson's disease I am describing, its development and changes that the disease brings along. Then I followed with the care of relatives person and support from the social system - the state.

In the practical part I have chosen for my research a survey, which was attended by only a small part of respondents from informal carers. Care of long-term sick family member is very difficult. Caregivers feel tired and exhausted, they report that their own health becomes worse. Care also affects caring professional life and there leisure time. Although half of the respondents are still able to find time for their hobbies. Some carers have no idea about the possibilities of helping services and just minimum of them use these services.