There is an urgent need to develop and/or expand palliative care for children in South Africa, and this editorial emphasises the scarcity of an evidence base on which to base clinical and operational decisions.

Children in Africa are more likely to face illness and death before the age of 5 years than anywhere else in the world. In South Africa, HIV infection is the leading cause of death among children aged less than 5 years, and there are approximately 300 000 children living with HIV/AIDS.

An essential tool in the spectrum of care offered to HIV-infected South African children is antiretroviral therapy (ART), and South Africa’s children’s ART programme is the largest in the world. Unfortunately, according to the national guidelines the programme currently reaches less than half of the children estimated to need ART. If the revised World Health Organization (WHO) guidelines for ART are considered, the proportion of children who need ART but are not accessing it is even greater.

Paediatric wards in South African government hospitals are occupied predominantly by children with HIV and AIDS-related illnesses. Providing hospital care for HIV-infected children is extremely stressful for both health care providers and caregivers, and sick children may face demoralising cycles of repeated hospital admissions. The needs of very ill young children are many and complex, and may be overlooked in busy, overstretched health care facilities.

Furthermore, not all sick children reach a health care facility and therefore are cared for in their own homes, placing a heavy burden on families, communities and informal cadres of health care workers, such as home-based care workers. In an attempt to broaden health care coverage the South African government has adopted a home-based care strategy premised on the belief that families are in the best position, with support from home-based care workers, to deliver a continuum of holistic care from infection through illness and death.

Holistic, ongoing care for very ill, dying and bereaved children and their families is situated within the emerging sub-specialty of paediatric palliative care, which focuses on achieving the best quality of life for children with life-threatening illnesses and their families.

Children’s palliative care is especially important in an African context because it can be delivered by a range of health care providers (professional and non-professional) and can be delivered both in health care facilities and in the home.

In South Africa, a number of health care facilities, hospices, non-governmental organisations, universities and the Department of Health are already delivering and developing aspects of children’s palliative care. The Hospice and Palliative Care Association of South Africa is affiliated to a wider African Palliative Care Association and has been active in children’s palliative care, particularly in advocacy and education.

Research elsewhere in Africa shows that when a nurse-led community-based children’s palliative care intervention is available there are increased referrals, increased prescriptions of essential drugs and improved compliance with treatment regimens. However, the findings from such studies cannot be directly transcribed to the South African context, where care of children is conducted by relatively unskilled home-based care workers, perhaps with little or no support from professionals trained in palliative care.

The marginalisation of palliative care research in Africa is well documented. Reasons given for this marginalisation include lack of skills and knowledge, professional isolation, poor patient accrual, high attrition, lack of agreement on outcome measures, and lack of a common language. There are also unique challenges in developing rigour in sampling design and in reporting. Importantly, there are ethical challenges to be considered when researching very sick children who are HIV infected.
Given the numbers of children affected and the need for palliative care to reach large numbers, I advocate that providing an evidence base for children's palliative care must be a priority. Any research should consider including perceptions and experiences of both caregivers and children, and researchers must consider cultural, language and ethical issues.

REFERENCES