different perspectives in Palestine. This paper reports on the perceptions of the physicians.

**Methods** This study used an ethnographic qualitative case study approach. It was conducted in one oncology unit in one Palestinian hospital and used two data collection methods: participant observation and semi-structured interviews.

**Results** The study generated 70 hours of observation and 35 interviews; physicians (n=5), nurses (n=11), children aged 6–18 (n=6), mothers (n=7) and grandmothers (n=6). The findings demonstrated that physicians were reasonably open with parents at the diagnosis stage; however, as the child’s condition got worse, they concealed negative information to protect parents from emotional suffering. However, despite their tendency to conceal information, they associated revealing information about the child’s deterioration to parents as an important act to protect themselves legally. Furthermore, they generally concealed negative information from children. There seemed to be a close link between the type, intensity and quality of information revealed and the stage of the cancer journey (diagnosis, treatment and prognosis).

**Conclusions** This study of Arab physicians delivering care to children with cancer and their families are positioned at a point in tension where they are reluctant to reveal information about a child’s deterioration and the knowledge that by concealing this information they place themselves in a difficult position. Although concealing information may be explained by cultural context, it does leave a communication gap for children and their parents at a very sensitive time in the child’s cancer journey.

**PUBLIC TRANSPORT AND HOSPICES – DO THEY LINK?**

Caroline Hart, Miriam Rigby, WM Cares. West Midlands Deanery

10.1136/bmjspcare-2019-ASP.28

**Background** Hospitals tend to be in central locations with good links to public transport including dedicated bus stops etc. This is due to very high numbers of both staff and patients needing 24 hour access. Hospices however tend to be less central, in quieter locations, making them less accessible to those who can’t drive; be they patients, their relatives or staff. So how easy are they to get to? And how easy is this information to find?

**Method** For all the hospices within the West Midlands with an inpatient unit (IPU); we looked at the following: what information was available on their website regarding access via public transport; how to access the IPUs via public transport using ‘traveline.org’ and Google. Finally, we contacted each hospice to ask if they had any written patient/visitor information regarding getting to the hospice.

**Results** We identified 12 hospices, and 15 IPUs in total. 5 had a link on their homepage indicating ‘how to find us’, 2 had information about nearest bus routes, 3 had information about the nearest train station. When contacting the hospices directly, only 3 hospices had written information available to give to patients and their relatives. We found that the distance from a bus stop to the hospice varied from a 2 to a 15 min walk, and more than half have no or very limited Sunday service.

**Conclusions** Most of the hospices in the West Midlands region have no information available regarding accessing the site using public transport, and limited transport services especially off peak.

Does this feed in to the inequalities in access to hospice services already seen across socioeconomic groups? In the first instance we hope this will encourage hospices to provide more travel information for patients, but could this also guide service development and future hospice location planning?

**THE AGE OF THE CARER HAS AN IMPACT ON THE BURDEN OF CARE EXPERIENCED BY THE CARER**

F Mukhtar, U Raja, S Singh, N Lovell, D Yi, U Higginson. King’s College London, Cicely Saunders Institute

10.1136/bmjspcare-2019-ASP.29

**Context** Those caring for people with chronic illnesses report high levels of unmet needs. Carers for patients with chronic illnesses must learn how to adapt with the demand of looking after a patient with a deteriorating condition. This can lead to increased stress due to sacrifices such as less time for socialising/financial difficulty.

**Objectives** To evaluate whether the age of the carer has impact on the burden experienced from caring for patients. Understanding the effect of caring for patients with chronic breathlessness and its burden would help interpret the results drawn from the main analysis of the OPTBreath study, which investigated the preference for the breathlessness support service.

**Methods** Answers to the carer burden inventory (CBI), included in the face to face interviews with carers were analysed by carer’s age. Categories of age were generated and answers to CBI items were converted into a total burden score, and synthesised by the scope review of literature on the carers’ wellbeing. relationships within the data and relating it to the results of other studies related to the wellbeing of carers. This includes journals and qualitative studies.

**Results** 65 carers completed 20 items in the CBI, The average age was 65 and 9 of 65 carers were men. 49/65 stated being both comfortable or coping on their current income, whereas only 10/65 stated in financial difficulty. Average burden score was 18.9: 12.5 in age <40; 16.4 in 40–49; 18.3 in 50–59; 20.7 in 60–69; 20.3 in 70–79 and 17.4 in 80+. The graphical illustration showed that there was an increasing trend from the <40 age range.

**Conclusions** Though the relationship was not linear the results indicate that there is clear larger burden of care on older carers. Suggestions as to why include increased morbidities in older carers and increased financial burden.

**CARERS’ REFLECTIONS ON PROVIDING INFORMAL END OF LIFE CARE AND THE POST-CARE PERIOD: A QUALITATIVE STUDY IN SOUTH GREATER MERSEYSIDE**

James Watson, Katherine Jehan. University of Liverpool

10.1136/bmjspcare-2019-ASP.30

**Background** End of life care need is increasing and will continue to do so. Government policy is for terminal patients to receive care and die in their preferred location. Most terminal patients prefer to receive care and die at home, but lack of palliative care funding places a substantial burden on informal carers. Inequalities exist in end of life care; informal carers