

a key protective factor in improved well-being. Differences between clinical areas were also identified.

Conclusion. There was a clear increase in distress calls and requests to prescribe or increase psychotropic medication to calm the distress during the 'lockdown'. Changes in behavioural presentation may have occurred partly due to the disruption to the complex systems that typically support a child and the shift away from community support. Children with intellectual disability and their families are unique and embedded in complex systems comprising schools, respite, and healthcare provision which work together to deliver optimal mental healthcare with psychosocial interventions with medication for higher-risk situations. Any shifts in these systems may lead to an imbalance and a higher likelihood of medication use.

Audit into post diagnostic support in newly diagnosed dementia patients

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Aims. This audit aims to identify whether newly diagnosed dementia patients are offered post diagnostic support and potential factors influencing patient choice.

Background. A diagnosis of dementia can be life changing and hence post-diagnostic support for dementia is key. Multiple guidelines suggest that post diagnostic support need to be offered to all patients diagnosed with dementia. The Department of Health and Social Care and other national/ local guidelines suggest that post diagnostic support is offered to all patients diagnosed with dementia.

Method. Data were collected for 40 patients diagnosed with dementia. Using random number generator, patient group was selected from pool of patients diagnosed with dementia between July' 2017 - December' 2017. Data included whether they had been offered support during the initial appointment and what post-diagnostic support was offered. Demographic details obtained to identify patterns of support accessed by patients.

Result. All patients were offered post-diagnostic support. Diagnosis was discussed in appointment in about 93% of patients. Medication was discussed in 82% patients. Driving was discussed in only 64% patients and LPA was discussed in only 63% patients. When given choice between Post diagnostic support group (PDSG) and Dementia adviser (DA), slightly more women tend to choose PDSG group. The only 2 ethnic minority patients chose DA. 21% more patients opted for PDSG group when they had a carer.

Conclusion. The positive is that some post-diagnostic support is offered to all patients. Although discussion of diagnosis with patients was done well, discussion of medication, driving and LPA can be improved upon. Ethnicity and family structure/ carer may have a bearing on patient choice of post-diagnostic support.

Medication charts and consent to treatment documentation audit in an acute mixed in-patient psychiatry unit in south Manchester

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Aims. The main aim of this audit was to look at documentation in medication charts in an acute mixed inpatient unit in South Manchester. In addition, we also looked at completion of capacity assessment and consent to treatment forms as appropriate.

Background. Safe prescription, administration and monitoring of medication is key to effective patient care. Due to the busy nature of inpatient hospital wards, errors do unfortunately occur both with the medications, and with the recording of their administration.

We will use a data collection tool to collect data as per standards described in our local GMMH policy. The medication chart will be used as the standard, as this is the current chart that is in use in the Trust.

Method. Data were collected from 31 medication charts for inpatients admitted in the ward between the 5/12/19 to 18/12/19. We captured data from each page of the medication chart that required a record to be made by any staff, including details of prescribing, administration and pharmacist checks. Data were recorded as either Yes/No or NA (Not Applicable). Data were then summarised and analysed using MS excel.

Result. Of the 31 patients, 22 (71%) had a capacity assessment form completed and 16 (52%) had a consent to treatment form completed. From the data analysis, it was clear that there are high rates of completion for the 'essential' parts of all prescriptions, including medicine name, dose, route and data. 'Route' was only recorded for 40% of prescriptions for depot medicines. Details of the administration of a medicine by a nurse was generally well-completed. For as required medications, all information relating to administration (date, time, dose and given by) were fully completed for 100% of prescriptions. For regular prescriptions however, the administration details were not as well-completed, where date of administration was recorded in 84% of prescriptions and signature in 29% of prescriptions. Unique patient identifiers are well-recorded on Page 1 of the prescription chart, though not maintained throughout the prescription chart. Nature of reaction to an allergy or sensitivity was only recorded in 6 of the 21 patients (29%).

Conclusion. Overall, there were good completion rates for the mandatory parts of the prescriptions. However improvements could be made for prescriptions as well as administration and pharmacy checks. The capacity assessment and consent to treatment forms could be improved upon too. We plan to put the recommendations and re-audit in 3-6 months' time.

Service evaluation and research project: pros and cons of centralisation of ECT services in Pennine Care NHS Foundation Trust

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Aims. The aim of this project would be to understand about the pros and cons/ benefits and risks of using a centralised model of ECT (currently being followed due to COVID restrictions) rather than a decentralised model of ECT (which has been the norm for a long time). A comparative account of both types of systems would help identify whether it could be implemented in Pennine Care and the results could be potentially transferable to other similar settings.

Method. An online survey was undertaken from staff members working in Pennine Care who have been involved in ECT delivery and management. The survey was undertaken between 07/12/