The impact of the Health and Social Care Act 2012 on forensic psychiatry

As a National Health Service forensic psychiatrist working on a newly commissioned low secure ward, the statement: ‘It is all too predictable that yet more patients will be pushed down forensic care pathways from which return to mainstream care will be difficult (p. 402)’ in Holloway’s excellent November editorial1 struck a firm chord with me.

In the past year, I have overseen an expansion of both the low secure forensic estate and the out of area patient placements. Although there was some clinical and commissioning intent to introduce the low secure estate to allow transition out of the medium secure estate (and indeed this has happened to some extent), there has been quite a surge of patients coming from the general acute services and the community.

We also receive some prison transfers; these include general adult community patients with no prior forensic history who were missed in the community owing to (poorly resourced) service lapses. Such patients become ‘forensic’ because of a lack of adequate community psychiatric services rather than being appropriate referrals to the service. In any case, we are expanding.

Good news for forensic staff, but not so good for patient care. Earlier psychiatric intervention for them may have even saved them from being locked up in prison. This is low-income country psychiatry in a high-income country.

At a recent presentation by some Californian psychiatrists, I was very impressed by the vigour with which they grapple with often very difficult legal circumstances of psychiatric care in their jurisdiction. They noted that most of their state hospital beds were occupied by their forensic patients. There was very little available for non-forensic patients, either in hospital or in the community. I wonder whether here in England we are also heading in that direction.

Finally, it appears that in this evolving, risk-focused, forensic-heavy psychiatric care environment, the ‘forensic’ patient today is not the same forensic patient from 20 years ago. These days, not every forensic patient is a high secure step-down patient. Why is it then more difficult to discharge forensic patients into the community, and return them to mainstream services? At the very least, the expanding low secure estate ought to provide an easier interface within the psychiatric services than was the case in the past. This way we will have done our best for our patients while contending with the difficult care environment being planned for us by this government. Indeed, who else will it?

Needless complexity in commissioning

Having attended a local third-sector and service user conference and having read the editorial by Holloway,1 I wonder whether the following needs more consideration.

It strikes me that dividing mental health commissioning responsibilities locally between the clinical commissioning groups (mental illness treatments) and local authorities (suicide and substance misuse prevention, mental health promotion) poses unnecessary complexity and bureaucratic waste. Despite lay representation in clinical commissioning groups, there is no democratic accountability similar to that offered through local councillors and local authority scrutiny committees which can call providers to attend a public meeting to account for their priorities in using public funds. Perhaps local elections might be more popular if electors realise that councillors could be voted out if they are not active in championing mental health issues such as dementia care. Furthermore, local authorities already have experienced procurement teams with ready access to performance management and audit functions.

Therefore, I wonder whether clinical commissioning groups should be relieved of all mental health commissioning responsibilities, with this function carried out entirely by local authorities. This would allow the commissioning groups to concentrate on acute and chronic medical diseases (which contribute to most of the cost via hospital bed usage and new technology). The added benefit of mental health being commissioned by local authorities would be integration of social and healthcare budgets for the benefit of people with severe mental illness such as psychosis and dementia. As a practising clinician, I find it difficult to separate social and health interventions in providing a good outcome for an individual patient; usually, there is a synergistic effect.

The other issue discussed by Holloway is ‘personalisation’. It is hoped that by April 2013, 70% of eligible mental health service users (mainly with severe chronic illness) will have a personal budget with an allocated broker to help clarify and achieve their choices in interventions. The above rationalising of commissioning would lend itself to a combined health and social care budget which can be spent pragmatically. A chip-and-pin charge card could be introduced to carry a combined budget, with greater accountability and freedom from having to collect receipts.

The third issue highlighted at the conference was an increasing body of evidence suggesting that active collaborations between statutory mental health providers and third-sector organisations result in better outcomes and lower number of bed days in psychiatric hospitals. Perhaps this should be considered an essential requirement for mental health trusts when submitting bids for a service.
Declaration of interest
These views are not those of my employer.


Prasanna N. de Silva. Consultant Old Age Psychiatrist, Tees, Esk and Wear Valleys NHS Foundation Trust, The Anchorage, Whitby, UK, email: prasanna@mentalhealthsolutions-whitby.co.uk
doi: 10.1192/pb.37.1.37a

Specialist community teams backed by years of quality research

In response to Dr Killaspy’s invited commentary1 on Dr Lodge’s piece favouring generic v. specialist mental health teams,2 professor Burns laments that ‘every change, no matter how hare-brained, is hailed “an innovation”’.2 He implies that it is ‘hare-brained’ to implement crisis response, early intervention and assertive community treatment (ACT) specialist teams, even though they all have unambiguously strong international evidence of both persistent effectiveness and economic advantage (e.g. Killaspy & Rosen,4 McCrone et al5).

We share Dr Lodge’s key concerns for continuity of care and the need to engage some individuals in long-term therapeutic relationships. For instance, ACT and early intervention psychosis (EIP) teams are specifically designed to amplify these functions, for those who need them and only while still needed. This has been readily addressed by having a generic front-end community mental health team (CMHT) co-located with primary care where possible and specialised back streams. This results in mutually supportive and often shared working between all these teams. Transfers, where they occur, are very slow, so continuity is preserved. Professor Burns and Dr Lodge argue from a false premise, as pitting generic against specialised teams is a ‘straw-man’ argument. They provide no evidence in support of retaining the generic status quo alone, just moral assertions. The status quo is often hailed as the ‘tried and tested’ condition to beat, when ‘there is surprisingly little evidence to show that [CMHTs alone] are an effective way of organising [community] services’, as stated in the National Institute for Health and Clinical Excellence guidance on managing schizophrenia in adults (CG82, p. 336).

Professor Burns accuses Dr Killaspy of being ungenerous, unjustified and disingenuous for standing up for systematised team approaches that have strong evidence internationally, in comparison with our more habitual comfort as clinicians with undifferentiated CMHTs and more traditional, hospital-centric and sedentary out-patient care. ‘Newer is not necessarily better’ he posits. Well, we appreciate his clinical conservatism. But, in stating that ‘Nobody waits to see if it makes any difference, never mind delivers an improvement’, how long does he wish us to wait, while depriving severely disabled UK citizens of an effective service delivery system (ACT) which has just been celebrated for more than 40 years since initial high-quality randomised controlled trials proved strongly favourable and cost-effective (e.g. studies by Stein, Test and Westwood), with waves of positive international replications since?

Over recent years, professor Burns and colleagues have muddied the waters by implying that indifferent results for even more diluted models of ‘intensive case management’ in the UK such as the UK700 and PRiSM studies somehow represented ACT, and proved that it did not provide any advantage in UK or Europe over CMHTs. They deem ACT to be unnecessary where, in comparison with other countries, there is an adequate health and Social Services ‘safety net’. Yet its effectiveness in Australia and Canada has been demonstrated in the context of a public health and welfare system at least as good as the UK’s at its best.4 Meanwhile, these much-vaunted ‘safety nets’ are now unravelling in many parts of Europe. This misleading position adopted by Burns and colleagues must bear some responsibility for this premature disinvestment, for the further dilution of these teams under financial pressure, and for the dampened enthusiasm for the UK research effort into ACT, when it has only just begun, with mixed results possibly owing to patchy team fidelity.4

Tragically, severely and persistently mentally ill Britons will suffer with neglect because of the partial dismantling or withdrawal of these essential integrative community care delivery systems. Community-based teams in the UK need their capacity to consistently follow the fidelity protocols of these specialist teams upgraded, not dismantled. This is a challenge to rigorous science, to sound commissioning, to communal action and ultimately to good government.

Declaration of interest
R.D. is editor of a consumer-oriented newsletter sponsored by Johnson & Johnson.

1 Killaspy H. Importance of specialisation in psychiatric services. Commentary on . . . How did we let it come to this? Psychiatrist 2012; 36: 364–5.


3 Burns T. Newer is not automatically better (e-letter). Psychiatrist 2012; 22 October.


A full list of references is available in an online version of this letter.

Alan Rosen. Professorial Fellow, School of Public Health, University of Wollongong, Brain and Mind Research Institute, University of Sydney, and Senior Consultant Psychiatrist, Far West Mental Health Service Sydney, Australia, email: alanrosen@med.usyd.edu.au. Leonard S. Stein. Emeritus Professor of Psychiatry, University of Wisconsin School of Medicine and Public Health, Madison, Wisconsin, USA. Patrick McGorry. Executive Director, OYH Research Centre, Professor of Youth Mental Health, University of Melbourne, Australia. Carol Harvey. Associate Professor, Department of Psychiatry, University of Melbourne, and North Western Mental Health, Melbourne, Australia. Max Birchwood, BPrA. Professor of Youth Mental Health, University of Birmingham, Clinical Director, YouthSpace Mental Health Service, Birmingham and Solihull Mental Health Foundation Trust, Birmingham, UK. Ron Diamond. Professor of Psychiatry, University of Wisconsin School of Medicine and Public Health, Madison, Wisconsin, USA.

doi: 10.1192/pb.37.1.38