A systematic review and synthesis of outcome domains for use within forensic services for people with intellectual disabilities†

Catrin Morrissey, Peter E. Langdon, Nicole Geach, Verity Chester, Michael Ferriter, William R. Lindsay, Jane McCarthy, John Devapriam, Dawn-Marie Walker, Conor Duggan and Regi Alexander

**Background**

There is limited empirical information on service-level outcome domains and indicators for the large number of people with intellectual disabilities being treated in forensic psychiatric hospitals.

**Aims**

This study identified and developed the domains that should be used to measure treatment outcomes for this population.

**Method**

A systematic review of the literature highlighted 60 studies which met eligibility criteria; they were synthesised using content analysis. The findings were refined within a consultation and consensus exercises with carers, patients and experts.

**Results**

The final framework encompassed three a priori superordinate domains: (a) effectiveness, (b) patient safety and (c) patient and carer experience. Within each of these, further sub-domains emerged from our systematic review and consultation exercises. These included severity of clinical symptoms, offending behaviours, reactive and restrictive interventions, quality of life and patient satisfaction.

**Conclusions**

To index recovery, services need to measure treatment outcomes using this framework.

**Declaration of interest**

None.

**Copyright and usage**

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Following de-institutionalisation, most people with intellectual disabilities live fairly independent lives in the community. There are 900,000 adults with intellectual disabilities in England, and estimates suggest that only around 3035 (0.3%) receive treatment in psychiatric hospital settings, with about half of them being in forensic hospitals.1–3 The health expenditure in this sector belies the low numbers, and it is estimated at over 300 million pounds sterling per annum.4,5 However, there is limited empirical information on service-level outcome domains and indicators, which in turn limits the ability to measure the effectiveness of these services. This is of concern in a health climate focused on outcomes6 and ‘payment by results’, but is even more relevant because of the recent government initiative to fundamentally transform care for people with intellectual disabilities.7 Although Fitzpatrick et al8 have conducted a systematic review of outcome measures used in generic forensic mental health services, and Gilbody et al9 completed a similar review of outcome studies in mental health, there has been no such work for forensic services providing care to people with intellectual disabilities. Further, although there has been a marked focus on recovery from mental health services, there has been comparatively little focus on this construct and its measurement within psychiatric hospital settings for people with intellectual disabilities, including forensic services. Recovery is often construed as ‘getting better’ or ‘reducing symptoms’, and within the context of in-patient services for people with intellectual disabilities, where there is often a focus on person-centred support and normalisation, including living as independently as possible within the community, the concept remains unclear, but the issues are not dissimilar from the ‘recovery’ debates within wider mental health services. However, recovery in the context of forensic services for people with intellectual disabilities, while subjective, should nevertheless incorporate the connectedness; hope and optimism about the future; identity; meaning in life; and empowerment (CHIME) framework,10 bearing in mind that some associated factors may be more proximal for this population (e.g. offending behaviours and stigma associated with disabilities).

In order to address these shortcomings, this study had the single aim of identifying the domains that should be used to measure outcome from forensic services for people with intellectual disabilities. Within the context of this project, outcome was defined as occurring at the level of the service as a whole, rather than individual outcomes associated with a specific treatment or intervention. In other words, we were primarily interested in outcomes that could index change over time across the entire range of interventions offered by a service, rather than outcome from a specific intervention (e.g. medication or psychological treatment), as this represents the real world of service delivery. Our aim was achieved within the context of two interrelated and iterative work streams: (a) undertaking a systematic review of studies that focused directly or indirectly on measuring outcomes from forensic services for people with intellectual disabilities and synthesising the findings into an initial framework of outcome domains, and (b) taking our initial framework and refining further within the context of a consultation exercise with patients and carers, as well as a two-round Delphi exercise with experts.

**Method**

An initial outcome framework was developed following a systematic review of the literature that focused on outcomes from forensic...
services for people with intellectual disabilities. As a starting point, and following discussion within the research team, we initially envisaged outcomes as falling into one of the three areas that were defined by the Department of Health as representative of quality. They are as follows: (a) effectiveness (e.g. the impact of generic treatment on health), (b) patient safety (e.g. untoward events as a result of treatment) and (c) patient experience of care (e.g. satisfaction).

Search strategy

The search strategy aimed to identify studies from a range of sources. Electronic databases searched on 1 June 2015 included Medline, Psyc (INFO), Embase, AMED, HMIC, BNI and CINAHL. Search terms employed were based on those used for a previous Cochrane reviews, for intellectual disability and forensic/offenders. The full search terms, including ‘explode’ terms, keywords and text words are included within our supplementary material. The systematic review is registered in advance with PROSPERO (registration number: CRD42015016941).

In order to ensure that no relevant publications were missed, the grey literature (opengrey.eu) was also searched using the keywords. The ancestry method was used to find suitable studies within the references of eligible papers. The ancestry method means searching the reference lists of papers that met our eligibility criteria for any further papers that may not have been previously included. In addition, expert members of the project team were consulted in order to identify any key references not retrieved by the search strategy as well as in press or unpublished articles.

Study selection and eligibility criteria

Duplicate studies were removed, and titles and abstracts of articles were screened against the eligibility criteria independently by two members of the research team (C.M. and N.G.). Any disagreements were resolved by a third reviewer (M.F.). Studies were included that (a) were published after 1980, as our initial searches revealed there was little relevant literature available before 1980; we opted to use this cut-off date to reduce the number of returned ineligible papers; (b) were in any language, as translations were obtained; (c) made use of any type of quantitative method; (d) involved adults within intellectual or autism spectrum disorders; (e) were older than 18 years of age; and (e) had current or past use of forensic services for people with intellectual disabilities, including community-based forensic services. Forensic services were defined according to the bed categories defined by the Royal College of Psychiatrists. This means that we included papers where the participants were either living within a high, medium or low secure in-patient forensic health service, or a forensic rehabilitation service, or they were living in the community, but receiving a service from a community-based forensic service. Studies were excluded if they only evaluated the effects of a specific intervention or treatment programme (e.g. randomised control trial of a medication or psychological treatment group), rather than examining outcomes at a service level.

Sixty studies met the inclusion criteria. None of the included studies were randomised controlled trials or meta-analyses. Twenty-eight studies were cohort outcome studies with follow-up from 1 to 20 years, and a further 32 were cross-sectional studies which reported service-level outcome data at one point in time. Several of these studies made use of the same or overlapping samples of participants, but as we did not make use of meta-analytic methods, this did not erroneously affect precision. The large majority of the studies included were from the UK, with only two studies originating elsewhere. Most studies made use of samples of men, as only two studies included women. Figure 1 depicts a flowchart outlining the study selection process and the number of studies identified at each stage.

Data extraction and analysis

Using a structured form, data were extracted from the included articles. Specifically, details regarding the sample, design, service type, methods, outcome domains and specific measures were obtained and coded. Content analysis was used to synthesise the outcome domains with reference to the three areas of quality as defined by the Department of Health, namely: (a) effectiveness, (b) patient safety and (c) patient experience. These three areas were used as an a priori superordinate framework. A process of refining and grouping similar outcome sub-domains together was then undertaken by two researchers. This led to the construction of a ‘framework’ to describe the outcome domains extracted from the eligible studies. This process is best described as both directed and summative content analysis because the process started with an a priori theoretical stance pertaining to service quality, followed by both coding and coding the extracted data, which was then interpreted within the context of our a priori theoretical stance. This methodology was advantageous because it allowed us to identify key concepts, consider their context and underlying meaning within and across studies, and develop coding variables, which were then refined into sub-domains.

Consultation exercise

Consultation groups

Following the completion of our systematic review, and the development of our initial outcome framework, we undertook three consultation groups with patients and one consultation group with carers to further consider and refine our outcome framework. Two of our patient groups took place within a high secure hospital in England, whereas the remaining groups took place within both a low secure hospital and a medium secure hospital, also in England. Participants were approached by the researchers, and the purpose of the group was explained using information sheets. Participants who agreed to take part provided informed consent. However, we were advised by our associated Research Governance office, within our National Health Service (NHS) Trust, that NHS Research Ethics opinion was not required for this project. Our groups included 3 women, 1 transgendered person and 11 men. For our consultation group involving carers, we recruited four participants from an existing carer group within a secure hospital, whereas two participants were recruited who were not part of this group. The carer participants had family members detained within three different secure hospitals.

Analysis

A semi-structured topic guide was used which was based around our initial outcome framework as a method to structure the conversations within our groups. Participants were encouraged to consider and discuss our initial framework, make modifications and choose outcomes they considered most important. The discussions were recorded and fully transcribed. The transcriptions were analysed using both directed and summative content analysis. As with our systematic review, this methodology allowed us to identify key concepts, context and meaning within transcripts, which were interpreted within our proposed framework. Any changes, or newly identified outcome sub-domains, were incorporated within our superordinate framework.

Delphi exercise

The Delphi method is an iterative and multi-staged structured process that can be used to develop group consensus. We made use of a two-round online Delphi exercise with expert clinicians, researchers and commissioners with experience of working within forensic services for people with intellectual disabilities.
Information about the study was advertised within the communication networks of existing stakeholder organisations within the UK (e.g. British Psychological Society). All participants were provided with information to help them make a decision as to whether they wished to take part in the study. Participants were presented with the revised outcome framework developed following our patient and carer consultation exercise. They were then invited to rate the importance of each sub-domain within each of the three superordinate domains along a 5-point Likert scale, where 1 was ‘not important’ and 5 was ‘extremely important’. Participants were also asked for their expert opinion about each sub-domain and whether they thought any additional outcome measures needed to be added. Finally, participants were asked to indicate the five sub-domains they considered to be the most important measures of outcome.

Following the completion of the first round, participants were invited to consider the responses of the group and re-consider their previous ratings. Those sub-domains with a mean rating of four or more were taken through to the second round, and participants re-rated their importance along the same 5-point Likert scale. Participants were also asked for their expert opinion about each sub-domain and whether they thought any additional outcome measures needed to be added. Finally, participants were asked to indicate the five sub-domains they considered to be the most important measures of outcome.

**Participants.** Seventeen participants took part in the first Delphi round, with 15 taking part in the second round. Nine participants were psychologists, seven were psychiatrists and one was a nurse. Participants were eligible to take part in the Delphi exercise if they were a clinician, researcher or commissioner with experience of working with forensic services for people with intellectual disabilities. Two participants identified themselves as having responsibility for commissioning, whereas a further two identified themselves as having both clinical and academic responsibilities.

**Results**

**Systematic review**

Using content analysis, data from eligible studies were extracted and categorised within the overarching superordinate domains: (a) effectiveness, (b) patient safety and (c) patient and carer experience. The complete list of identified sub-domains that emerged following our analysis, along with the associated studies, is found in Tables 1–6. For simplicity, studies have been divided into cohort, retrospective cohort, cross-sectional or case study designs. These findings were synthesised into our initial framework of outcomes which was taken forward and used within our consensus exercises (Table 7).
| Study and setting | Design | N | Outcome sub-domain | Measure/indicator |
|------------------|--------|---|--------------------|-------------------|
| **Alexander et al.**<sup>17</sup> Medium secure | Retrospective cohort study Follow-up: up to 13 years | 64 | Treatment response/outcome | Level of CPA support, MHA status and CGIS |
| | | | Discharge pathway | Level security/type of placement at discharge |
| | | | Readmissions | % of patients readmitted to the same unit |
| | | | Relapse in MH | % of patients who experienced a relapse in MH symptoms |
| | | | Reoffending | Number of patients formally reconvicted, cautioned or had police contact |
| | | | ‘Offending-like’ behaviour | Behaviour which could be classed as an offence but did not lead to police contact |
| **Alexander et al.**<sup>18</sup> Medium secure | Retrospective cohort study Discharged patients Follow-up: 6 years | 138 | Length of stay | Mean and median number of days |
| | | | Discharge outcome | % of positive (move to a lower level of security) or negative (move to higher level of security) discharges |
| | | | Discharge pathway | Level security/type of placement at discharge |
| **Alexander et al.**<sup>17</sup> Medium secure | Retrospective cohort study Follow-up: 6 years | 138 | Length of stay | Mean and median number of days |
| | | | Discharge outcome | % discharged successfully (to a lower level of security) or not (to same/higher level of security) |
| **Alexander et al.**<sup>19</sup> Medium secure | Retrospective cohort study Follow-up: up to 6 years | 30 | Length of stay | Discharged patients only, mean and median days |
| | | | Discharge outcome | % of discharged patients who moved to either a lower or same/higher level of security |
| **Alexander et al.**<sup>20</sup> Medium secure | Retrospective cohort study (Discharged patients. PD v. ID v. PD+ID) Follow-up: up to 9 years | 145 | Length of stay | Median number of days and age on discharge |
| | | | Clinical symptoms | PCL:SV pre-treatment scores (clinician rated) |
| | | | Risk assessment | HCR-20 pre-treatment scores (clinician rated) |
| | | | Reoffending | Reconvictions at 1, 2 and 5 years follow-up. Categorised as serious/violent offending (as defined by Home Office) |
| | | | Time to reoffend | The difference between discharge date and reconviction date for subsequent offence (median) |
| **Ayres & Roy**<sup>21</sup> Community | Case series (70% male) Follow-up: up to 3 years | 26 | Cost | Average saving cost per patient (based on hourly rate of support) and for the service over 3 years |
| | | | Level of support | Hours per annum providing both direct/indirect support |
| **Benton & Roy**<sup>22</sup> Community | Cohort study Follow-up: 9.5 months average | 61 | Clinical symptoms | ABC and Mini PAS-ADD (clinician rated) |
| | | | Treatment engagement | Categorised treatment use and% with history of service use |
| | | | Reoffending | Number of patients who reoffended; mean number of offences per patient |
| | | | ‘Offending-like’ behaviour | Telephone call with carers regarding police contact |
| **Benton & Roy**<sup>23</sup> Community | Cohort study Follow-up: up to 3 years | 113 | Reoffending | Number of patients arrested, reconvicted and cases dropped |
| | | | Discharge pathway | Level security/type of placement at discharge |
| | | | Discharge outcome | % of referrals discharged |
| | | | Prevention of in-patient admissions | Number of patients whose in-patient admission was prevented via community treatment |
| **Butwell et al.**<sup>24</sup> High secure | Cohort study Follow-up: 10 years Up to 278 | Length of stay | Median and mean years, calculated per episode |
| | | | Discharge pathway | Level security/type of placement at discharge |
| **Day**<sup>25</sup> Medium secure | Cohort study Follow-up: 3 years | 20 | Length of stay | Mean number of months |
| | | | Treatment response | Categorised as either good (settled and co-operative), fair (continuing lesser problems) or poor (severe problems) |
| | | | Discharge pathway | % of patients discharged to rehab villa, community or hostel |
| | | | Discharge outcome | Level of adjustment at follow-up, based on personal knowledge, hospital notes and liaison with involved agencies |
| | | | Readmission | % of patients readmitted to the same unit |
| | | | Reoffending | Number who reoffended or returned to prison |
| Study and setting | Design | N | Outcome sub-domain | Measure/indicator |
|------------------|--------|---|--------------------|-------------------|
| **Dickens et al**<sup>26</sup> Medium and low secure Follow-up: 15 months | Retrospective cohort study | 48 | Length of stay, Clinical symptoms | Mean number of days, HONOS-Secure: change between baseline and final rating. Quartile points used to allow for different lengths of stay (clinician rated) |
| **Fitzgerald et al**<sup>27</sup> Medium secure Follow-up: 2 years | Retrospective cohort study | 145 | Reoffending, Risk assessment | Home Office records; % of patients with both general and violent offences, OGRS |
| **Gray et al**<sup>28</sup> Medium secure Follow-up: 2 years | Retrospective cohort study | 115 | Reoffending, Risk assessment | Home Office records; % of patients with both general and violent offences, HCR-20, VRAG, PCL:SV (rated once) |
| **Halstead et al**<sup>29</sup> Medium secure Discharged patients Follow-up: up to 13 years | Cohort study | 35 | Length of stay, Treatment outcome | Mean and median number of days, rated as either good (risks reduced, safe for discharge), some (progress made in some areas, risk remains), none (no change) or poor (worse than admission), % of patients discharged, Level security/type of placement at discharge and follow-up, Recurrence of symptoms of illness or challenging behaviour, % of patients readmitted to hospital, Behaviour which could be interpreted as an offence, Number of patients readmitted |
| **Lindsay et al**<sup>30</sup> Community Follow-up: 13 years | Cohort study | 62 | Offending-like behaviour | Percentage of patients 'suspected of' reoffending |
| **Lindsay et al**<sup>31</sup> Community Follow-up: 7 years | Cohort study | 184 | Reoffending, Harm reduction | Number of patients with clear evidence of reoffending, Decrease in the number of incidents 2 years before referral vs. incident data at follow-up |
| **Lindsay et al**<sup>32</sup> Community Females only Follow-up: 3 years | Case series | 18 | Reoffending | Number of patients with clear evidence of reoffending |
| **Lindsay et al**<sup>33</sup> Community Follow-up: 12 years | Cohort study | 247 | Reoffending, Harm reduction | Number of patients with clear evidence of reoffending, Decrease in the number of incidents 2 years before referral vs. incident data at follow-up |
| **Lindsay et al**<sup>34</sup> Community Follow-up: up to 20 years | Cohort study | 309 | Reoffending, Harm reduction | Number of patients with clear evidence of reoffending, Standardised service incident report data; number of offences committed in the 2 years before referral vs. follow-up reoffending data |
| **Lindsay et al**<sup>35</sup> Mixed services Follow-up: 2 years | Cohort study | 197 | Discharge pathway | Number of patients discharged and level security/type of placement at discharge, 1-year and 2-year follow-up, Local agency data and arrest frequencies used, Number of patients who completed treatment |
| **Linhorst et al**<sup>36</sup> USA Community service Follow-up: 6 months | Cohort study | 252 | Reoffending, Treatment response/engagement | % of patients who reoffended (even if this did not lead to police contact) determined by interview with current care team, Mean and median number of years, % of patients in high, medium, low and community services |
| **Marks**<sup>37</sup> [unpublished thesis] Medium secure Follow-up: 4 years | Retrospective cohort study | 28 | Reoffending, Length of stay, Discharge pathway | % of patients who reoffended (even if this did not lead to police contact) determined by interview with current care team, Mean and median number of years, % of patients in high, medium, low and community services |
| **Morrissey & Taylor**<sup>38</sup> High secure Follow-up: 2 years | Case reports | 13 | Clinical symptoms, Discharge outcome | YSQ (patient rated), IPDE and PCL:SV (clinician rated); change in mean scores, Number of patients remaining in treatment, discharged to medium secure or transferred to another high secure ward |
| **Morrissey et al**<sup>39</sup> High secure Follow-up: 5 years | Retrospective cohort study (ID v. other non-ID services) | 70 | Risk assessment | HCR-20: change in mean scores |
| **Morrissey et al**<sup>40</sup> High secure Follow-up: 2 years | Prospective cohort study | 73 | Discharge outcome | % of patients who made positive move to a lower level of security, % of patients who made positive move to a lower level of security or negative (the same level of security) progress |
| Study and setting | Design | N          | Outcome sub-domain                     | Measure/indicator                                      |
|-------------------|--------|------------|----------------------------------------|--------------------------------------------------------|
| Morrissey et al  | Retrospective cohort study (Two cohorts: in-treatment and admission cohort) | 68 and 24 | Length of stay, Clinical symptoms, Discharge outcome, Incidents | HCR-20, PCL-R (clinician rated) |
| High secure       | Follow-up: up to 6 years              |           |                                        |                                                        |
|                   |                                   |           |                                        |                                                        |
| Palucka et al     | Retrospective cohort study          | 20        | Length of stay, Discharge outcome, Discharge pathway, Incidents | Median number of days, Number of patients discharged, Level security/type of placement at discharge, Any criminal justice involvement (even if charges were not pressed) |
| Canadian in-patient unit | Follow-up: up to 9 years            |           |                                        |                                                        |
| Reed et al        | Retrospective cohort study          | 45        | Length of stay, Discharge pathway, Discharge outcome, Incidents | Mean number of weeks, Level security/type of placement at discharge, Positive (discharged to a level of lower security) or negative (discharged to a level of higher security) outcome |
| Low secure        | Follow-up: up to 14 years           |           |                                        |                                                        |
| Xenitidis et al   | Retrospective cohort study          | 64        | Length of stay, Discharge pathway, Discharge outcome, Incidents | Mean number of months, Level security/type of placement at discharge, Good (discharged to community) or bad (not placed in a community setting) outcome, Collected from hospital incident records. Incidents at baseline (week 6 to 10 of stay) were compared to end of stay (last 4 weeks of treatment). Frequency (total number of incidents per month) was adjusted for length of stay. Change in incidents calculated per person per week |
| Low secure        | Follow-up: up to 11 years           |           |                                        |                                                        |

ABC, Aberrant Behaviour Checklist; Mini PAS-ADD, Mini Psychiatric Assessment Schedules for Adults with Developmental Disabilities; YSQ, Young Schema Questionnaire; IPDE, International Personality Disorder Examination; PCL-R, Psychopathy Checklist – Revised; EPS-BRS, Emotional Problems Scale – behaviour rating scale; EPS-SRS, Emotional Problems Scale – self report scale; CPA, Care Programme Approach; MHA, Mental Health Act; MH, mental health; CGIS, Clinical Global Impressions Scale; PCL:SV, Psychopathy Checklist: Screening Version; HCR-20, Historical, Clinical, Risk Management-20; HoNOS, Health of the Nation Outcome Scale; OGRS, Offender Group Reconviction Scale; VRAG, Violence Risk Appraisal Guide.

a. The sample in Alexander et al overlaps with that of Alexander et al and Esan et al.
b. The sample in Fitzgerald et al overlaps with that of Gray et al.
c. The sample in Lindsay et al overlaps with that of Lindsay et al.
d. The sample in Reed et al overlaps with that of Xenitidis et al.
## Table 2: Summary of studies presenting data on the outcome domain of effectiveness (cross-sectional studies)

| Study and setting | Design | N   | Outcome sub-domain | Measure/indicator |
|-------------------|--------|-----|---------------------|-------------------|
| Ajmal et al. | High secure | Cross-sectional | 79 | Clinical symptoms | GSI and RSES (patient rated) |
| Beer et al. | Low secure | Cross-sectional | 59 | Placement appropriateness | Percentage of patients assessed as requiring a less secure placement |
| Beer et al. | Low secure | Cross-sectional | 68 | Length of stay | Mean number of months |
| Chaplin et al. | Low secure | Cross-sectional | 22 | Risk assessment | HCR-20 median scores |
| Chilvers & Thomas | Medium secure | Cross-sectional | 77 | Clinical symptoms | NAS-Pi scores (patient rated) |
| Crossland et al. | High, medium and low secure | Cross-sectional | 60 | Length of stay | Median number of months |
| Dickens et al. | Medium and low secure | Cross-sectional | 68 | Incidents | Severity rated by the individual completing the form as either: near miss, minor, moderate, high or very high |
| Esan et al. | Medium and low secure | Cross-sectional | 114 | Length of stay | Mean and median months for both discharged and in-treatment patients |
| Fitzgerald et al. | Medium and low secure | Cross-sectional | 136 | Incidents | Number of patients involved in incident in 6-month period |
| Hall et al. | Medium and low secure | Cross-sectional | 136 | Treatment needs | Clinician ratings |
| Hogue et al. | High, medium, low and community | Cross-sectional | 228 | Clinical symptoms | EPS-BRS (clinician rated) |
| Johnson | Medium and low secure | Cross-sectional | 44 | Clinical symptoms | RSES and EBS (patient rated) |
| Kellett et al. | High secure | Cross-sectional | 45 | Clinical symptoms | BSI (patient rated) |
| Lindsay et al. | Community | Cross-sectional | 52 | Offender-like behaviour | Percentage of patients suspected of reoffending |
| Lindsay et al. | High, medium and low secure | Cross-sectional | 212 | Risk assessment | HCR-20, VRAG, Static-99, SDRS, RM-2000 |
| Lindsay et al. | High, medium, low secure and community | Cross-sectional | 197 | Risk assessment | EPS-BRS (clinician rated) |
| Study and setting | Design | N  | Outcome sub-domain | Measure/indicator |
|-------------------|--------|----|--------------------|-------------------|
| Lothhouse et al\(^6\) | Cross-sectional 5 months of data | 64 | • Length of stay  
• Risk assessment  
• Incidents | • Mean number of years  
• CuRV  
• Aggression defined as acts of physical violence, aggression, force to hurt or damage to staff, peers or environment. Included verbal abuse which was aggressive, threatening or caused offence. Two researchers rated each incident as: ‘agression present’ or ‘agression absent’. Number of patients who were aggressive in month 1 versus month 5 |
| Mansell et al\(^7\) | Cross-sectional NHS versus private provider units | 1891 | • Delayed discharge  
• Incidents | • Percentage of patients who had completed treatment but did not have any plans to leave the service in the next month  
• Average frequency where a patient was hurt by a patient or staff member (per patient over a 6-month period) |
| McMillan et al\(^8\) | Cross-sectional 6-month period | 124 | • Risk assessment  
• Incidents | • MDT ratings per patient on risk of physical violence (scale of 0-8) and number of times patient had been violent in 6 months prior to risk assessment  
• Author coded each description based on explicit criteria and guidelines. e.g. physical violence (attempted, contact between assailant or object and victim, evidence of physical harm to victim or attendance of medical personnel). Coded from computerised hospital database |
| Morrissey et al\(^9\) | Cross-sectional 12-month period | 60 | • Incidents | • Coded as either interpersonal physical aggression or verbal aggression/aggression to property. Further rated as low, medium or high risk of harm. Number of patients involved in an aggressive incident  
• HCR-20  
• PCL-R and EPS-BRS (clinician rated) |
| O’Shea et al\(^10\) | Cross-sectional 3-month period following | 109 | • Risk assessment  
• Incidents | • Hospital records in 3-month period following risk assessment for aggression and self-harm. Coded using Overt Aggression Scale. Rated on severity (1-4). Number of patients involved in any incident |
| Perera et al\(^11\) | Cross-sectional 15-month period of incidents | 388 | • Length of stay  
• Delayed discharge | • Median number of years and percentage of patients who had stayed longer than 5 years  
• Percentage of patients assessed as requiring a less secure placement |
| Thomas et al\(^12\) | Cross-sectional | 102 | • Length of stay  
• Delayed discharge  
• Security need  
• Treatment needs | • Mean and median number of years  
• Percentage of patients assessed as requiring a less secure placement and main reason for this  
• CANFOR-Short and CANDID-Short. Average number of needs and unmet needs |
| Uppal & McMurrant\(^13\) | Cross-sectional (ID sample included in wider data-set) 15-month period of incidents | 396 | • Incidents | • Hospital computerised reporting system. Coded as per Department of Health: Category A: major incidents (e.g. abscond, hostage taking); Category B: serious incidents (e.g. serious assault involving a weapon, attempted suicide); Category C: untoward incidents (e.g. attempted abscond, assault without a weapon); Category D: all other incidents (minor assault and verbal abuse)  
Most frequent location and time of incident  
Percentage of incidents which were violent and which were self-harm  
Average monthly figure generated |

GSi, Global Severity Index; RSES, Rosenberg Self Esteem Scale; SBS, Social Behavioural Schedule; MOAS, Modified Overt Aggression Scale; NASS-P, Novaco Anger Scale and Provocation Inventory; ASD, autistic spectrum disorder; VRAG, Violence Risk Appraisal Guide; RSES, Rosenberg Self Esteem Scale; EBS, Evaluative Beliefs Scale; BSI, Brief Symptom Inventory; SDTS, Security, Dependency and Treatment Needs Scale; CANFOR, Camberwell Assessment of Need for Adults with Developmental and Intellectual Disabilities; ID, intellectual disability.
Table 3 Summary of studies presenting data on the outcome domain of patient safety (cohort studies)

| Study and setting               | Design                        | N    | Outcome sub-domain                                      | Measure/indicator                                                                 |
|--------------------------------|-------------------------------|------|---------------------------------------------------------|------------------------------------------------------------------------------------|
| Alexander et al19               | Retrospective cohort study    | 30   | Seclusion, restraint and intensive observations          | Mean number of episodes per patient/month                                          |
| Medium and low secure           | Follow-up: 6 years            |      |                                                         |                                                                                    |
| Alexander et al18               | Retrospective cohort study    | 138  | Seclusion, restraint and intensive observations          | Mean number of episodes per patient/month (adjusted for length of stay)           |
| Medium secure                   | Follow-up: 6 years            |      |                                                         |                                                                                    |
| Alexander et al17               | Retrospective cohort study    | 138  | Seclusion, restraint and intensive observations          | Mean and median episodes per patient/month                                         |
| Medium secure                   | Follow-up: 4 years            |      |                                                         |                                                                                    |
| Ayres & Roy21                   | Case series                   | 26   | Medication                                              | Case study: reduction in frequency of use of pro re nata medication               |
| Community                       | Follow-up: up to 3 years      |      |                                                         |                                                                                    |
| Butwell et al24                 | Cohort study                  | Up to 278 |                                             | Death                                                                             |
| High secure                     | Follow-up: 10 years           |      |                                                         | Frequency of episodes (n and % of all patients)                                    |
| Morrissey & Taylor22            | Cohort study                  | 13   | Seclusion                                               | Hours per patient for every 6 months of treatment                                  |
| Medium and low secure           | 6-month period                |      |                                                         |                                                                                    |
| Mason23                         | Retrospective cohort study    | 64   | Seclusion                                               | Episodes at baseline (week 6 to 10 of treatment) compared to end of stay (last 4 weeks of treatment) |
| Low secure                      | Follow-up: up to 11 years     |      |                                                         |                                                                                    |

Table 4 Summary of studies presenting data on the outcome domain of patient safety (cross-sectional studies)

| Study and setting               | Design                        | N    | Outcome sub-domain                                      | Measure/indicator                                                                 |
|--------------------------------|-------------------------------|------|---------------------------------------------------------|------------------------------------------------------------------------------------|
| Esan et al24                    | Cross-sectional (ASD v. Non-ASD) | 114  | PRN usage, Restraint, seclusion and intensive observations | Mean number of episodes (total frequency divided by total number of months of stay to provide an average monthly figure) |
| Medium and low secure           | 6-month period                |      |                                                         |                                                                                    |
| Mansell et al21                 | Cross-sectional              | 1891 | PRN usage, Seclusion, restraint, locked areas, Access to healthcare | Average number of episodes                                                        |
| Medium and low secure           | 12-month period               |      |                                                         | Percentage of units who reported a delay in patients accessing primary (nurse/dentist) healthcare |
| Mason22                         | Cross-sectional              | 36   | Seclusion                                               | Number of patients secluded; average number of seclusion episodes per patient/year; reason for seclusion and distress-related behaviours after seclusion |
| High secure                     | 12-month period               |      |                                                         |                                                                                    |

Table 5 Summary of studies presenting data on the outcome domain of patient and carer experience (cohort studies)

| Study and setting               | Design                        | N    | Outcome sub-domain                                      | Measure/indicator                                                                 |
|--------------------------------|-------------------------------|------|---------------------------------------------------------|------------------------------------------------------------------------------------|
| Fish & Lobley26                 | Cohort study (4)              | 20   | Quality of life                                         | QoLS: change from pre- to post-move                                             |
| Community                       | Follow-up: 1 year             |      |                                                         |                                                                                    |
| Long et al23                    | Cohort study (4)              | 10   | Milieu                                                  | EssenCES change from pre to post move                                            |
| Low secure                      | Follow-up: 3 months           |      |                                                         | In-patient satisfaction questionnaire: change from pre to post move               |
| Female only                     |                               |      |                                                         |                                                                                    |
| Marks20 (unpublished thesis)    | Retrospective cohort study    | 28   | Quality of life                                         | QoLS scores during treatment                                                     |
| Medium secure                   | (4) Follow-up: 4 years        |      |                                                         |                                                                                    |
| Trout20 (unpublished report)    | Cohort study (4)              | 44   | Quality of life                                         | PWI scores: change from pre to post move                                          |
| High secure                     | Follow-up: up to 2 years      |      |                                                         | Service specific evaluation questionnaire using a visual Likert scale via interview: change from pre to post move |

QoLS, Quality of Life Scale; EssenCES, Essen Climate Evaluation Scale; PWI, Personal Wellbeing Index.
Effectiveness

Fifty-three studies were categorised as presenting data that involved at least a single outcome that attempted to measure effectiveness (Tables 1 and 2). Our analysis led to 12 sub-domains within the effectiveness superordinate domain (Table 7). These included sub-domains such as length of stay, discharge outcome, clinical symptoms, treatment responsiveness, reoffending behaviours and risk assessment.

As a sub-domain, length of stay was considered within 22 studies (Tables 1 and 2), and varied between 1 and 9 years across included studies. However, it was recognised that as a measure of outcome, length of stay is problematic because (a) it tended to be reported for only those who had actually been discharged, rather than the entire in-patient population, and (b) it is complicated because some patients move from one hospital to another, and data may not capture their entire length of stay across all hospitals. As another sub-domain, discharge outcome was considered within 16 studies (Tables 1 and 2) and was defined as moving to an increasing or decreasing level of security within or across forensic hospitals, or discharge to a community-based setting. Several of the included studies focused on delayed discharge and highlighted the difficulties with finding appropriate accommodation that mitigated risk.

Sixteen studies were judged to have included sub-domains that were classified as falling within the clinical symptom sub-domain and these are detailed in Tables 1 and 2. These included measures that made use of clinician or patient ratings of clinical symptomatology. However, only two studies reported change in clinical symptoms over time for a cohort of patients. A variety of tools were used to index change over time within this sub-domain and included such measures as the Brief Symptom Inventory, Emotional Problem Scales, Mini Psychiatric Assessment Schedules for Adults with Developmental Disabilities (mini PAS-ADD), Health of the Nation Outcome Scale (HoNOS)-Secure, and Clinical Global Impressions Scale. Treatment responsiveness was also coded as a sub-domain, but it was recognised that this is intertwined with the clinical symptom sub-domain; this was included as a separate sub-domain because it focused on whether a patient was likely to be responsive to treatment efforts, rather than the actual response.

Reoffending and risk were classed as separate sub-domains, with 18 and 12 studies considering variables within these sub-domains, respectively (Tables 1 and 2). Most commonly, studies tended to focus on reoffending using data derived from police or Ministry of Justice records. One study followed up reoffending at 1, 2 and 5 years post-discharge, whereas another set of studies using the same data-set reported on whether there was reoffending behaviour within the 2 years following discharge from hospital. Another, based in Australia, considered arrest data and ‘any criminal justice involvement’ following discharge. A series of studies, based in the community, considered whether treatment within the context of a community-based forensic service led to a reduction in offending behaviours. It is important to note that many people with intellectual disabilities may not be formally dealt with by criminal justice agencies, and as a consequence, formal arrest and conviction data may not be a valid index of reoffending. Hence, the category of ‘reoffending-like behaviour’ described in these studies is one which is important because it is likely to have increased validity.

Risk also emerged as a likely sub-domain which could be used to index the effectiveness of forensic services for people with

### Table 6: Summary of studies presenting data on the outcome domain of patient and carer experience (cross-sectional studies)

| Study and setting | Design | N | Outcome sub-domain | Measure/indicator |
|-------------------|--------|---|--------------------|------------------|
| Langton et al. | Cross-sectional | 18 | Milieu | CIES read aloud to patients, scores during treatment |
| Mansell et al. | Cross-sectional | 118 | Service satisfaction/complaints | Number of patient generated complaints per unit over 6-month period, recorded via standardised survey |
| Steptoe et al. | Cross-sectional | 28 | Quality of life | SOS and LEC scores during treatment |
| Willetts et al. (2014) | Cross-sectional | 45 | Milieu | EssenCES scores during treatment |

CIES, Correctional Institutions Environment Scale; SOS, Significant Others Scale; LEC, Life experience checklist.
intellectual disabilities. The vast majority of these studies reported on their use of structured clinical judgement tools, such as the Historical, Clinical, Risk Management-20 (HCR-20),\textsuperscript{20} with several others considering actuarial risk assessment measures, such as the VRAG,\textsuperscript{28,53,54} RM-2000 or STATIC-99.\textsuperscript{56,59,60} There was only a single study that considered how changes in scores on a risk assessment tool may relate to treatment outcome from forensic services for people with intellectual disabilities.\textsuperscript{57}

Patient safety

Eleven studies were categorised as presenting data that were considered to index outcome within the patient safety domain. Premature death was considered by one study, which differentiated between suicide and death associated with natural causes, whereas another study incorporated physical health, an important and relevant sub-domain considering the high rates of morbidity amongst forensic populations, including people with intellectual disabilities.

The five sub-domains that emerged following our analysis also included those related to ‘reactive’ or ‘restrictive’ interventions such as the use of physical interventions and seclusion, \textit{pro re nata} (PRN) medication or a change in observations levels. ‘Reactive’ or ‘restrictive’ interventions fall within the safety domain defined by the Department of Health.\textsuperscript{8,81} We have adopted the same approach here.

However, there is an overlap with the previously discussed effectiveness domain. ‘Reactive’ or ‘restrictive’ interventions can be construed as proxy variables for behaviour, and their use may correlate with increasing behaviour difficulties. However, they are not an intervention and instead are reactive strategies taken to try to manage behaviour difficulties in the short term to ensure safety. However, medical, psychological and social care interventions developed using a formulation that aim to rehabilitate and/or habilitate are not ‘reactive’, and as such, these would fall within the effectiveness domain. This includes psychological and social interventions, as well as medication prescribed to treat a diagnosed mental illness or distressing symptoms.

Patient experience

Within this superordinate domain, 11 studies were categorised as capturing outcomes related to patient experience which were categorised into 4 sub-domains. These were quality of life, therapeutic milieu, patient involvement and patient satisfaction. Four studies included in the review measured quality of life using a number of ratings scales, such as the Quality of Life Questionnaire\textsuperscript{86} or Life Experience Checklist,\textsuperscript{71} whereas three other studies focused on therapeutic milieu or ward atmosphere using either the Correctional Institutions Environment Scale\textsuperscript{70} or the EssenCES Climate Evaluation Scale.\textsuperscript{59,72} Three studies focused on patient satisfaction in response to service development, and only a single study considered patient involvement as an indicator of outcome.

Consultation exercise

Following the completion of our systematic review, and the development of our initial outcome framework (Table 7), this was presented to our consultation groups with patients and carers. Revisions were made and the revised outcome framework was used within our Delphi exercise with experts. As with the systematic review, we made use of the three superordinate domains (a) effectiveness, (b) patient safety and (c) patient and carer experience as a framework for our analysis of the data generated from our consultation exercises.

Consultation groups

**Effectiveness.** Several patients expressed the view that length of stay should be an important index of outcome; several said they were frustrated because they thought that length of stay was excessive for many patients. However, some carers expressed an alternative view, stating that a shorter length of stay may be problematic and lead to premature discharge. Patients from high security settings were of the opinion that discharge to medium security was indicative of positive progress, whereas for those in medium and low security, discharge to a community-based service was seen as positive. Several carers further discussed how frequent moves between hospitals and wards can be particularly destabilising and may actually be associated with a negative outcome.

The appropriateness of a placement with respect to meeting treatment needs was discussed and considered important by many carers and patients. One stated, ‘I would much rather be further away for eight to nine months [and get the right treatment] than be nearer for 18 months’. Another commented, ‘it is very important for people to go to a place where they are happy, not just because it is closer to family’.

Many commented further about the importance of much needed clinical interventions being available within each service, focusing specifically on psychological treatments and appropriate levels of meaningful activity. Carers spoke about wanting and needing individually tailored care pathways focusing on patient need rather than rigidly designed care pathways that were not based upon a formulation of treatment needs. One said, ‘it has got to be individually led’, and another commented, ‘he needed an individualised package of support which was right for him’. Alongside this, carers also expressed the view that a similar individualised package of support needed to be made available to patients when discharged into the community, with one stating, ‘I worry about the fact that the service wasn’t there in the community…there is so little support in the community’.

Improvements in clinical symptoms and behaviour were recognised by both patients and carers as indicative of positive change. This included quantifiable changes in the frequency of incidents, including improvements in communication and a reduction in angry feelings. Different patients stated, ‘before I wouldn’t engage in conversation and now I’ve learnt different strategies so I don’t kick off so often’, and ’a reduction in incidents, reduction in restraints, using diversion more frequently, preempting incidents’. Carers broadened this by commenting that some patients may not fully understand what they need to achieve to move forward, as there is often too much focus on measuring incidents within services. Some patients also shared this view and stated, ‘there is too much focus on incidents and not on understanding them … taking back to the beginning of the process as opposed to just dealing with what the consequences are’. Carers considered that a family member may be able to make a more nuanced judgement about changes to clinical symptoms because of their long-standing knowledge of the patient. Both patients and carers commented on the importance of engagement with services and therapies as positive indicators of progress within this sub-domain.

Both patients and carers agreed that ‘staying safe’ once discharged was a positive outcome, recognising that a reduction in risk was associated with a positive outcome, and several carers adopted the position of both carer and potential victim, expressing concern that their own safety could be compromised. One carer said, ‘if they said take him home I would be too scared’.

Finally, within the effectiveness superordinate domain, and as an addition to our initial framework, adaptive functioning
was considered by patients to be an important indicator of outcome. They talked positively about how they hoped that staying in hospital would bring about improvements in adaptive functioning, such as budgeting, occupational skills and broader life skills. One said, ‘I have been given skills like cooking and cleaning…’. However, several commented that staying in hospital may be associated with a loss of adaptive functioning, and several said they thought they have lost skills. For example, one said, ‘other hospitals let patients get real jobs. I want this to happen in this hospital’, while another commented, ‘since we’ve been locked up here we don’t get a chance to do that sort of thing [budgeting] so you don’t know what to do when you get your money’. One of our carer participants commented, ‘he used to be able to do things. He’s lost those skills since he’s been here’.

**Patient safety.** Patients spoke about how a reduction in aggression and the use of seclusion was a relevant outcome measure, as did a number of carers, whereas patients also spoke about being victimised by other patients in hospital. Some carers further considered that taking medication regularly was an indicator of positive outcome, and they also spoke about how a planned reduction in medication could also be a positive outcome. For example, one commented, ‘if he could come off olanzapine, that would be progress’, whereas another stated, ‘a reduction in PRN medication and other medication is a goal’.

Several carers and patients expressed concern about polypharmacy and side-effects, indicating that they felt this was a restrictive practice and alluding to the possibility that medication may be used to sedate in order to control behaviour; one commented, ‘he’s never been on this amount of medication … he’s so heavily dosed up … if he’s been medicated to manage his behaviour, he’s not learned how to manage his behaviour’. One of the patients strengthened this view by commenting, ‘can you be careful about medication and patients being overdosed’.

**Patient and carer experience.** This superordinate domain was modified as a consequence of our consultation groups in order to incorporate carer experience, alongside the experiences of service users. Carers spoke about whether they were satisfied with the level of care being afforded by their family member and indicated that this was an important measure of outcome. Several spoke about being satisfied with the care being offered by the hospital. One stated, ‘it is a dream come true; the place where he is now, it’s lovely… it’s a dream for places like that to be about’, whereas another commented, ‘the hospital are [sic] fantastic; the staff are fantastic and at long last somebody is realising the amount of problems he has got and that is one of the problems I had before’. Others considered the importance of having a sense of security as a consequence of the quality and responsiveness of care being given to their relative; one stated, ‘not having to be worried about him; if we died tomorrow, services would be there for him and do what was best for him without thinking of the cost’. However, several carers spoke about having to fight or battle for service provision and felt that sometimes services did not listen or involve them appropriately in the care pathway. This was illustrated by the following, ‘there was nothing we could say which would be taken on board … it was very much ‘no’ this is what we think’, and ‘I was asking for help in the community for years before my son was admitted to hospital’.

Patients and carers considered the importance of quality of life as an indicator of positive outcome, and many spoke about having hopes for a job, relationships and involvement in their local communities, with well-integrated high-quality support. Several carers emphasised the importance of high-quality accommodation once a patient was discharged, and one commented, ‘he would be in accommodation that was specifically designed for people with autism, but he had sufficient support with people who actually understood his condition and were able to spot the warning signs so I didn’t have to keep flagging them up’. Another stated, ‘he needs an individual planned package with sufficient staff and appropriate training’. Carers also commented that leaving hospital was not the end of patients’ journeys and spoke about the importance of continuing to monitor outcome and progress over the longer term, rather than view the ‘case as closed’. Others spoke about valuing having in-patient services which could be used in times of crisis; this was illustrated by the following comment, ‘… for him to go back to a secure unit because he’s a danger when he does deteriorate’.

The availability of and engagement with meaningful activity was seen as a potential indicator of positive outcome by both patients and carers. They spoke about having employment, and how increasing engagement in activities could be indicative of improvement. Carers spoke further about the importance of having meaningful activities available within hospital settings and went on to further consider how developing and maintaining social networks are further evidence of a positive outcome. This included developing and maintaining positive relationships within family, friends, and pets, and further included romantic relationships.

**Changes to our initial outcome framework.** A variety of changes to our initial outcome framework were made following the analysis of the data from our consultation groups. This included the incorporation of additional sub-domains or the modification of sub-domains. Specifically, we changed the label of the superordinate domain ‘patient experience’ to ‘patient and carer experience’. Considering the effectiveness superordinate domain, we made changes as follows: (a) treatment response and recovery and clinical symptom severity were modified to include carer ratings of clinical improvement, (b) acquiring adaptive skills was added as a new sub-domain, as was (c) engagement with therapies and services. Within the patient safety superordinate domain, we incorporated (a) safeguarding and victimisation, as a new sub-domain, whereas (b) overuse of medication was strengthened by making reference to unacceptable side-effects and patient satisfaction with prescribed medication. Finally, within the patient and carer experience superordinate domain, we incorporated sub-domains focusing on (a) the carer experience incorporating both communication and involvement, (b) closeness to home area and (c) the level of support and involvement within the community, as well as access to occupational activities. We also included that quality of life could be indexed by either clinicians or the patient.

**Delphi exercise**

Following our revisions to the outcome framework, we completed a two-round Delphi exercise with experts in order to create consensus about the most important outcomes for forensic services for people with intellectual disabilities. None of the sub-domains were rated as ‘not important’ or ‘slightly important’ by the participants. Five sub-domains did not reach consensus at the end of round one, and these were (a) length of stay, (b) security needs, (c) adaptive functioning, (d) clinician-rated quality of life and (e) closeness to home area. Participants were asked to rate five outcomes that they thought were the most important, and length of stay was included within these top five and was therefore retained and taken through to round two.

Six sub-domains received the highest average ratings by experts at the end of round two where a clear consensus emerged. These were (a) discharge outcome, (b) treatment response/engagement,
(c) premature death and suicide, (d) therapeutic milieu, (e) meaningful activity and (f) reoffending/offending-like behaviour. However, when asked to indicate their top five sub-domains, participants chose sub-domains exclusively within the effectiveness superordinate domain, and these were (a) clinical symptom severity/treatment needs, (b) reoffending/offending-like behaviour, (c) treatment response/engagement/insight, (d) risk assessment measures and (e) recovery measures/direction of care pathway. Perhaps this is not surprising, considering that all of the participants had current or past clinical responsibility for patients within services.

Integrating the findings, the final most important sub-domains were (a) discharge outcome, (b) recovery measures/direction of care pathway, (c) treatment response/engagement/insight, (d) clinical symptom severity, (e) reoffending/offending-like behaviour, (f) risk assessment, (g) premature death and suicide, (h) therapeutic milieu and (i) access to work and meaningful activity. The findings from the Delphi exercise were considered and synthesised into our findings from the consensus exercises and our systematic review. This led to the emergence of a final outcome framework. These are important indicators of the quality of forensic services, but may not always directly relate to clinical effectiveness.

**Table 8** Final framework of outcome domains and sub-domains

| Source of domain | Stage 1 | systematic review; Stage 2, patient/carer involvement groups; Stage 3 = Delphi, CSJ, criminal justice system, PPA, pro re nata, BNF, British National Formulary. |
|------------------|---------|-----------------------------------------------------------------------------------------------------------------------------------------|
| **Effectiveness**|         |                                                                                                                                         |
| Discharge outcome/direction of care pathway | 1 |                                                                                                                                         |
| Delayed discharge/current placement appropriateness | 1 |                                                                                                                                         |
| Readmission (i.e. readmitted to hospital or prison) | 1 |                                                                                                                                         |
| Length of hospital stay | 1 |                                                                                                                                         |
| Adaptive functioning | 1 |                                                                                                                                         |
| Clinical symptom severity/treatment needs: patient rated | 1 |                                                                                                                                         |
| Clinical symptom severity/treatment needs: clinician rated | 1 |                                                                                                                                         |
| Recovery/engagement/progess on treatment goals: clinician rated | 1 |                                                                                                                                         |
| Recovery/engagement/progess on treatment goals: patient/carer rated | 2 |                                                                                                                                         |
| Reoffending (i.e. charges/convictions) on discharge | 1 |                                                                                                                                         |
| Offending-like behaviour (no CSJ involvement) on discharge | 1 |                                                                                                                                         |
| Incidents (violence/self-harm) (in care setting) | 1 |                                                                                                                                         |
| Risk assessment measures | 1 |                                                                                                                                         |
| Security need (i.e. physical/procedural/escort/leave) | 1 |                                                                                                                                         |
| **Patient safety** |         |                                                                                                                                         |
| Premature death/suicide | 1 |                                                                                                                                         |
| Physical health | 1 |                                                                                                                                         |
| Medication (i.e. PRN usage/exceeding BNF limits/side-effects) | 1/2 |                                                                                                                                         |
| Restrictive practices (restriction) | 1 |                                                                                                                                         |
| Restrictive practices (seclusion/segregation) | 1 |                                                                                                                                         |
| Victimization/safeguarding | 1 |                                                                                                                                         |
| **Patient and carer experience** |         |                                                                                                                                         |
| Patient experience: involvement in care | 2 |                                                                                                                                         |
| Patient experience: satisfaction/complaints | 1 |                                                                                                                                         |
| Quality of life: patient rated | 1 |                                                                                                                                         |
| Therapeutic climate | 1 |                                                                                                                                         |
| Access to work/meaningful activity (where appropriate) | 2 |                                                                                                                                         |
| Level of support/involvement in community (post discharge) | 2 |                                                                                                                                         |
| Carer experience: communication with services/involvement in care | 2 |                                                                                                                                         |

The aim of this project was to identify the domains that should be used to measure outcome for people with intellectual disabilities and forensic needs. This is a topic relevant to all psychiatrists, particularly following the abuse scandal at a specialist intellectual disability hospital in England, Winterbourne View, and the resulting agenda to care for people with intellectual disabilities within ‘mainstream’ psychiatric services.18 Similar issues are of concern around the world as many work toward the social inclusion of people with intellectual disabilities within mainstream services within the health and social care sectors. Our aim was achieved by undertaking a systematic review coupled with a consultation exercise involving patients, carers and experts. The findings revealed a series of important sub-domains spread across three superordinate domains indicative of quality.22 These captured a range of clinical and patient safety variables, along with factors measuring both the patient and carer experience of care.

The largest outcome domain was effectiveness, which is not surprising. The sub-domains included were those that captured aspects of the care pathway, along with a focus on clinical symptoms, recovery and a reduction in reoffending. Related variables, such as length of stay, discharge and need for security, were included, but these may not always directly correlate with clinical need. For example, it would be possible for someone who has received successful treatment to remain in hospital due to delayed discharge because of difficulties with the provision of community-based services to manage risk. Further, length of stay in this context, as an indicator of outcome, should be neither too short nor too long, and instead should be ‘just right’ as it should be appropriate to meet the needs of individual patients, adding substantial complexity, especially when considered as a sole indicator of outcome. As such, focusing on multiple sub-domains allows for a richer and more thorough picture of the circumstances surrounding the care being offered to patients within forensic services.

However, consideration as to what ‘effective treatment’ in this context actually looks like requires further exploration, both on an individual patient level and on a wider service level. Only one study5 described the nature of the treatment programme that is delivered within the service. Effective treatment is likely to form a combination of appropriate medical, psychological and social intervention, informed by individual clinical formulations, but the availability is likely to vary across services, depending on patient needs. At present, in deciding whether a service is effective, regulatory and commissioning bodies rely on easily measurable process variables (e.g. the existence or otherwise of various policies, and the availability or otherwise of various treatments) rather than paying attention to the more important question of whether any of this is making a difference to the outcome. This is clearly unsatisfactory. Likewise, there are no studies which have looked at the economic evaluation of treatments, a rather surprising finding considering the abundance of anecdote and opinion in this field about costs.5 Considering the future, the structure and form of ‘effective treatment’ within forensic services should be clarified and drawn from a robust evidence base, bearing in mind that there are very few clinical trials to identify the most effective intervention across the range of those that are available. As such, greater investment in research investigating the clinical effectiveness of forensic services for people with intellectual disabilities is needed.

There are other sub-domains clustered around safety and the patient and carer experience which we incorporated into our final framework. These are important indicators of the quality of forensic services, but may not always directly relate to clinical effectiveness. Nevertheless, helping to ensure that patients with intellectual disabilities detained in forensic services have a good quality of life,
including access to meaningful activity, is a core business of such services. Measuring sub-domains with the broader safety and patient and carer experience domain is clearly important, for all stakeholders, especially patients and their carers. The measurement strategy for these domains could be standardised nationally, bearing in mind that they may not correlate directly with clinical effectiveness.

Related to this, the addition of a series of consultation exercises, alongside our systematic review, adds a particular strength to our project. This helped to ensure that we adequately captured the views of all stakeholders and appropriately synthesised them into our final framework. It is important to note that although experts tended to focus on clinical outcomes, patients and carers tended to focus more upon the quality of service provision, and the experience of receiving a service, alongside clinical outcomes. As such, it became important to ensure that these findings formed part of our final outcome framework.

Contrasting our framework with that developed by Fitzpatrick et al., there are both similarities and differences. Fitzpatrick et al. grouped outcome measures across a variety of similar domains, such as recidivism, service outcomes, mental state, compliance, satisfaction and substance misuse, among others. They were able to successfully review a variety of specific outcome measures that would enable measurement across these domains, whereas in our study there are relatively fewer instruments that have been standardised for use with people with intellectual disabilities across the sub-domains we have included within our framework. At the same time, there were some noted differences between our framework and that reported by Fitzpatrick et al. For example, substance misuse did not feature explicitly within our framework, but nevertheless is an issue for many with intellectual disabilities, and would fall easily within our Incidents sub-domain. Conversely, there were specific sub-domains that we included which did not appear within the framework reported by Fitzpatrick et al., such as adaptive functioning, access to meaningful activity, as well as the use of restrictive practices, which no doubt are all issues for those with forensic mental health problems and are likely to be more salient with services for people with intellectual disabilities.

**Clinical implications**

The findings from the current project have direct relevance to recent government initiatives, including Building the Right Support and the new National Service Model that were developed and published in response to the institutional abuse that took place at Winterbourne View in England. For many years, there has been a focus on ensuring that people with intellectual disabilities are afforded good quality care within their own communities, rather than in hospital, and the abuse that occurred at Winterbourne View has reignited the drive to ensure that people with intellectual disabilities are afforded good quality care within their own communities, alongside active, clear and robust discharge planning. In order to achieve these aims, services need to be able to measure outcomes, and for those who are admitted to in-patient forensic services, it would be valuable for care and treatment reviews to be structured around our outcome framework. This would help ensure that decisions about care are based on the research evidence and on indicators that are considered to measure change appropriately, helping to ensure the process is robust. One of the further important findings from our work is that we have integrated the findings from the evidence base which was used as the springboard to develop our framework. Although there are difficulties with many of the included studies, what was apparent was the absence of a focus on recovery and exploration of the subjective meaning of recovery in this context. Alongside this, many of the studies were small and very few longitudinal studies drawing on a well-developed outcomes framework have been completed, which is both clearly and sorely needed.

**Limitations**

All of these recommendations need to be balanced against several weaknesses associated with this study. First, our findings from the systematic review are based on the research evidence. Inherently, our findings from the systematic review are only as robust as the quality of the research that was reviewed. The predominant issue with many of the studies that were included was that few were longitudinal studies measuring outcomes, demonstrating that these outcomes had validity and reliability as an outcome indicator. Related to this, because of the marked variation across studies in terms of methodology, it became impossible to find a suitably reliable and valid tool that would index quality in this context. Moreover, if we had been able to measure study quality, this would not have altered the weight put on one study as opposed to another, because the focus was on the domains and how those domains were measured.

Although this is problematic, it is attenuated by the consultation exercises with patients, carers and experts. The patients included within our focus groups are vulnerable, detained under the Mental Health Act, and are often not given a voice. They directly contributed to the development of our outcomes, telling us what was important to them, as users of the services. Our findings from the consultation exercises were incorporated into our methods which helped to ensure that our findings were shaped carefully by those affected by our findings, which in turn increased validity. The second weakness is that the findings from our consensus exercises are based upon the views of a group of individuals, and only four carers were included; our findings may have been enhanced with a larger number of carers, but the content had become repetitive suggesting we had reached content saturation. Although we attempted to capture the views of a variety of patients, carers and experts, it is certainly possible that had we asked a different group of patients, carers and experts, different issues may have emerged from our analysis. Third, as our study included participants from the UK, there is a question as to whether the findings are generalisable to healthcare systems in other countries. However, we would anticipate that the findings have implications within other countries offering similar services and could be used to inform further research within similar hospitals and services in other parts of the world.

Finally, and looking forward to the future, further work is needed to investigate the reliability and validity of our outcomes framework. This may lead to a reduction in the number of sub-domains found within our current outcomes framework, which would increase the probability that services would integrate the framework into their
services. Related to this, it is important to consider that individual-level outcomes are likely to be very important when indexing recovery, and further work is needed as to how these are measured across services, because they are likely to be associated with local clinical practices which may be idiosyncratic and vary from service to service. However, a degree of standardisation would be valuable when monitoring and improving service outcomes, and specifying the method of measurement across our sub-domains is an important next step. Together, our framework should have a beneficial impact on improving both service quality and patient outcomes, whereas it would also allow for the creation of a national minimum data-set, specific to these services, which could be used to track patient outcomes and help develop and refine care pathways. Considering future research, it is now appropriate to consider the likely instruments that could be used to measure outcomes, allowing us to trial this framework within existing hospital care pathways.

Catrin Morrissey, PhD, Division of Psychiatry and Applied Psychology, School of Medicine, University of Nottingham, Nottingham, and Complexity and Forensic Service, Lincolnshire Partnership NHS Foundation Trust, Lincoln, UK; Peter E. Langdon, PhD, Tizard Centre, University of Kent, Canterbury, and Broadland Clinic, Hertfordshire Partnership University NHS Foundation Trust in Norfolk, Norwich, UK; Nicole Geach, Milles, Division of Psychiatry and Applied Psychology, School of Medicine, University of Nottingham, Nottingham, UK; Verity Chester, MSc, Department of Psychiatry, Partnerships in Care, Norfolk, and Norwich Medical School, University of East Anglia, Norwich, UK; Michael Ferriter, PhD, [Retired from] Forensic Division, Nottinghamshire Healthcare NHS Trust, Nottingham, UK; William R. Lindsay, PhD, Department of Psychology, University of the West of Scotland, UK, and Department of Psychology, The Danshell Group, UK; Jane McCarthy, FRCPsych, Department of Forensic and Neurodevelopmental Sciences (F-ANS), Institute of Psychiatry, Psychology & Neuroscience, King’s College London, London, UK; John Devapriam, FRCPsych, Department of Psychiatry, Leicestershire Partnership NHS Trust, Leicester, and Care Quality Commission, London, UK; Dawn-Marie Walker, PhD, Health Sciences, University of Southampton, Southampton, UK; Conor Duggan, OBE, FRCPsych, Institute of Mental Health, University of Nottingham, Nottingham, UK; Regi Alexander, FRCPsych, Department of Psychiatry, Partnerships in Care, Department of Psychiatry, Leicester Partnership NHS Trust, and Department of Psychiatry, University of Leicester, Leicester, UK.

Correspondence: Regi Alexander, Partnerships in Care, Lion Road, Palgrave, Dixs, Norfolk, IP22 1BA. Email: regialexander@nhs.net

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