RESEARCH PAPER

Intensive upper limb neurorehabilitation in chronic stroke: outcomes from the Queen Square programme

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ABSTRACT

Objective Persistent difficulty in using the upper limb remains a major contributor to physical disability post-stroke. There is a nihilistic view about what clinically relevant changes are possible after the early post-stroke phase. The Queen Square Upper Limb Neuromechanism programme delivers high-quality, high-dose, high-intensity upper limb neurorehabilitation during a 3-week (90 hours) programme. Here, we report clinical changes made by the chronic stroke patients treated on the programme, factors that might predict responsiveness to therapy and the relationship between changes in impairment and activity.

Methods Upper limb impairment and activity were assessed on admission, discharge, 6 weeks and 6 months after treatment, with modified upper limb Fugl-Meyer (FM-UL, max-54), Action Research Arm Test (ARAT, max-57) and Chedoke Arm and Hand Activity Inventory (CAHAi, max-91). Patient-reported outcome measures were recorded with the Arm Activity Measure (ArMA) parts A (0–32) and B (0–52), where lower scores are better.

Results 224 patients (median time post-stroke 18 months) completed the 6-month programme. Median scores on admission were as follows: FM-UL = 26 (IQR 16–37), ARAT = 18 (IQR 7–33), CAHAi = 40 (28–55), ArmA-A = 8 (5–12) and ArmA-B = 38 (24–46). The median scores 6 months after the programme were as follows: FM-UL = 37 (IQR 24–48), ARAT = 27 (IQR 12–45), CAHAi = 52 (IQR 35–77), ArmA-A = 3 (IQR 1–6.5) and ArmA-B = 19 (IQR 8.5–32). We found no predictors of treatment response beyond admission scores.

Conclusion With intensive upper limb rehabilitation, chronic stroke patients can change by clinically important differences in measures of impairment and activity. Crucially, clinical gains continued during the 6-month follow-up period.

INTRODUCTION

Stroke remains common1 and persistent difficulty in using the upper limb is a major contributor to ongoing physical disability.2 The general consensus remains that most spontaneous recovery of the upper limb occurs over the first 3 months after stroke and current levels of rehabilitation result in little improvement after that, particularly at the level of impairment.3 Improving outcomes through higher dose (time in rehabilitation or number of repetitions) and intensity (dose per session) of rehabilitation is an attractive option.4 However, clinical trials of higher dose upper limb rehabilitation have generally produced the magnitude of improvement that will change clinical practice,5 whether delivered in the early6 or chronic stages post-stroke.7,8 A common factor in these trials is that the dose (in hours) of additional therapy remained relatively low (18–36 hours). Despite scepticism that stroke patients could tolerate much higher doses,9 one study managed to deliver 300 hours of upper limb therapy to chronic stroke patients over 12 weeks and reported changes in measures of both impairment and activity that were far greater than those in lower dose studies.10 Three hundred hours represents an order of magnitude higher than any dose of rehabilitation offered in previous upper limb rehabilitation trials and deserves further consideration. However, this idea is challenging because of the logistics of setting up such a trial in healthcare settings where the ethic of high-dose, high-intensity rehabilitation is not supported. In this context, it is important to report the findings of clinical services that are able to deliver higher doses than conventionally seen. The Queen Square Upper Limb (QSUL) Neurorehabilitation programme is a single-centre clinical service that provides 90 hours of timetabled treatment focusing on the post-stroke upper limb in chronic (>6 months post-stroke) stroke patients. Here, we report (i) outcomes for patients admitted to this programme at the National Hospital for Neurology and Neurosurgery, University College London Hospitals NHS Trust (UCLH), including 6-month follow-up data to look at whether any clinical benefits were maintained, (ii) the characteristics of the patients admitted and any predictors of response and (iii) the relationship between changes in impairment and activity.

METHODS

Participants

Patients were referred by primary care physicians for advice about ongoing management of the paretic upper limb following stroke. There were no criteria around time since stroke. Clinical criteria warranting admission to the programme were broad but were focused on whether we felt able to help patients achieve their goals for the affected upper limb. Barriers to admission included factors precluding the ability to work in upper limb tasks: (i) absent movements, throughout the limb; (ii) a painful shoulder limiting an active forward reach (mostly due to adhesive capsulitis); (iii) severe spasticity or non-neural loss of range and (iv) unstable medical conditions. These patients were not offered admission to the treatment programme but were provided with appropriate advice or referred for...
other treatment approaches. Re-referral to the programme was then considered (eg, after successful management of adhesive capsulitis of the shoulder). Features that were more likely to result in admission to the programme included at least some ability to flex the shoulder and at least visible flickers of movement in finger and/or wrist extensors, although this was not always the case as reflected in baseline upper limb scores. Patients were admitted to the programme as day attenders, either from home or from the UCLH-dedicated patient accommodation and as such were either self-caring or self-caring with the support of one other person.

Interventions
Initial assessment consisted of analysis of both movement and performance in activities of daily living. Subsequent treatment was aimed at reducing impairment and promoting re-education of motor control within activities of daily living. Individualised meaningful tasks were practiced repeatedly in order to facilitate task mastery with a focus on quality of movement. This was achieved through (i) adaptation of the task, for example, decomposing tasks into individual components to be practiced; (ii) adaptation of the environment, for example, fabrication of functional splints and adaptation of tools such as cutlery or screwdrivers, to enable integration of the affected hand in meaningful activities; (iii) assistance, for example, de-weighting the arm to allow strengthening and training of movement quality and control through increased range and (iv) independent task practice. Coaching was considered a key component of the programme and used throughout to embed new skills and knowledge into individual daily routines. Consequently, individuals increase participation and confidence in their desired goals, enhancing self-efficacy and motivation to sustain behavioural change beyond the end of the active treatment period.

This overall approach was achieved through two daily sessions each of physiotherapy and occupational therapy, supplemented with tailored, individualised interventions, including repetitive practice with a rehabilitation assistant or robotic device, sensory retraining, use of dynamic and functional orthoses, neuro-muscular electrical stimulation and group work. Furthermore, patients were encouraged to work on cardiovascular fitness during the programme. A 6-hour timetable was implemented 5 days a week for 3 weeks (total therapy time, 90 hours). Motor tasks could be described as passive or active, assisted or unassisted, and functional or non-functional. Over the 3 weeks, the aim was to increase the time spent on active, unassisted functional tasks, depending on a patient’s level of impairment and progress. The programme was staffed with a 1:1 staff to patient ratio (two physiotherapists, two occupational therapists and two rehabilitation assistants for six patients).

Quantitative assessment
All patients had the following baseline scores measured on admission (table 1): modified Rankin scale (mRS), Barthel Index (BI), Neurological Fatigue Index (NFI), Hospital Anxiety and Depression Scale (HADS) and sensation (as indexed by light touch on palm as described in Fugl-Meyer, FM, assessment of the upper extremity).

The affected upper limb was assessed on admission (T1), discharge (T2), 6 weeks (T3) and 6 months (T4) postdischarge, using the following measures: FM (upper limb) (FM-UL), Action Research Arm Test (ARAT), Chedoke Arm and Hand Activity Inventory (CAHAI-13) and Arm Activity Measure (ArmA).

The FM-UL is a stroke-specific, performance-based impairment index (where impairment refers to loss of body structure and function) with good validity and intra-rater and inter-rater reliability.11,12 Here, we have used a modified version of the FM-UL, excluding measures of coordination and reflexes based on hierarchical properties of the scale,13 and as these do not relate to the upper limb synergies of interest.14 The minimum clinically important difference (MCID) has been reported as 5.25 points.15

The ARAT assesses a patient’s ability to handle objects differing in size, weight and shape and therefore can be considered to be an arm-specific measure of activity limitation.16 The MCID has been suggested as 5.7 points.17

The CAHAI-13 is a validated, upper limb measure that uses a 13-point quantitative scale in order to assess recovery of the arm and hand in performing activities of daily living after a stroke.18 It is a performance test using day-to-day items. It is not designed to measure the patient’s ability to complete the task using only their unaffected hand, but rather to encourage bilateral upper limb use. No MCID has been reported for CAHAI-13 (although the minimum detectable change is reported as 6.218).

The ArmA is a patient-reported outcome score19 with two components. The ArmA-A asks patients about their ability to care for their affected arm either themselves with their unaffected arm or by a carer or a combination of both of these. ArmA-B asks patients about how easy or hard it is to use the affected arm in activities of daily living. For ArmA-A and ArmA-B, note that lower scores are better. No MCID has been reported for ArmA-A or ArmA-B.

Analysis plan
Our primary aim was to measure changes in upper limb deficit over time which were examined using the Friedman test for all outcome scores. Post-hoc analysis to test differences between individual time points was carried out with the Wilcoxon signed-rank test. As a secondary aim, we were then interested in whether baseline characteristics correlated with final (T4) outcomes. We initially examined for correlations between individual baseline characteristics and outcomes using Spearman’s rank correlation. We also tested whether median T4 outcomes were different depending on gender or side of lesion using Wilcoxon rank sum test. Time since stroke was not considered as a covariate as once in the chronic phase, there is no indication that time since stroke has a linear effect on outcome. Lastly, we performed multiple linear regression to look for predictors of either absolute scores at T4 or the change in scores T1–T4. Lastly, we also examined the relationship between the changes in outcome scores from admission to 6-month follow-up using Spearman’s rank correlation.

RESULTS
Baseline characteristics
Between January 2015 and December 2017, 268 patients were admitted to the QSUL programme (representing 46% of patients assessed in out-patient clinic). 40.4% of patients were admitted from home while 59.6% stayed in the UCLH-dedicated patient accommodation. Note, only 16 patients were admitted to the programme prior to 6 months post-stroke, and only one patient prior to 3 months, reflecting overall referral patterns to the programme.

Of those, 30 were non-stroke (12 traumatic brain injury, 8 brain tumour, 3 peripheral neurological conditions, 3 spinal cord injury and 4 inflammatory central nervous system conditions)
changes in upper limb deficit

The median scores for the affected upper limb at admission (T1), discharge (T2), 6 weeks (T3) and 6 months postdischarge (T4) are shown in table 2. The Friedman test was used to demonstrate a significant effect of time since admission for all outcome scores; FM-UL ($\chi^2 (3)=431.8, p<0.0001$); ARAT

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Upper limb scores before and after admission</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Admission</td>
</tr>
<tr>
<td>ARAT</td>
<td>18 (7–33)</td>
</tr>
<tr>
<td>CAHAI</td>
<td>40 (28–55)</td>
</tr>
<tr>
<td>ArmA-A</td>
<td>8 (4.5–12)</td>
</tr>
</tbody>
</table>

All scores given as median (IQR).
(χ² (3)=383.2, p<0.0001); CAHAI (χ² (3)=371.6, p<0.0001); ArmA-A (χ² (3)=238.4, p<0.0001) and ArmA-B (χ² (3)=305.6, p<0.0001). Post-hoc analysis with Wilcoxon signed-rank test was conducted to test the hypothesis that the difference between the paired values (T1−T2, T1−T3, T1−T4, T2−T3, T2−T4, T3−T4) comes from a distribution whose median is zero (ie, testing whether the change in score is significantly greater than zero). A Bonferroni correction was applied to the significance level for the six comparisons made for each of the five different outcome scores (p<0.05/30=0.0017). For each outcome score, there was a significant difference between each pair of time points compared, except T2−T3 for the ArmA-A (online supplementary table). Individual scores at each time point are shown in figures 1 and 2. Individual changes in scores in comparison to admission are shown in table 3 and figures 1 and 2. These results demonstrate improvement during the 3-week programme but also continued improvement after discharge.

By 6 months after the programme, 68.3% of patients had achieved greater than the MCID of 5.25 points on the FM-UL. For ARAT, this figure was 61.6%. No published MCID is published for the other scores. MCID is often quoted as 10% of the maximum score. If this were the case, then by 6 months, admission (eg, FM1 if attempting to predict FM4 or change from FM1 to FM4). Models predicting T4 scores for FM-UL, ARAT, CAHAI, ArmA-A and ArmA-B were also highly significant (p<0.0001). Models predicting change from T1 to T4 for ARAT, CAHAI, ArmA-A and ArmA-B were all highly significant (p<0.0001). However, the only factor that not differ by gender or affected hemisphere. There was a trend for patients with nondominant affected limbs to achieve better (lower) ArmA-B scores in comparison to those with dominant affected limbs (median 17 for nondominant vs 27 for dominant, p=0.027, uncorrected). There were no differences in T4 scores between patients with dominant or nondominant affected limbs for any of the other outcome scores.

Second, we were interested to see whether any baseline characteristics (T1) correlated with magnitude of change from admission to 6-months follow-up (T1−T4). Spearman's rank correlation was used to show that FM-UL, ARAT, CAHAI and ArmA-B at T1 correlated with change in ARAT and CAHAI (but not FM-UL) from T1 to T4. FM-UL at T1 also correlated with change in both CAHAI and ArmA-A from T1 to T4. ArmA-B at T1 correlated with change in both ArmA-A and ArmA-B from T1 to T4. Age, NFI, HADS, BI and mRS at T1 did not correlate with any scores at T4. All p values were corrected for multiple comparisons by Bonferroni correction (table 5). We also performed a Wilcoxon ranked sum test to show that change in scores from T1 to T4 was not different depending on gender or affected hemisphere. There was a trend for patients with nondominant affected limbs to improve on the ArmA-B between T1 and T4 to a greater degree than those with dominant affected limbs (median changes of −17.5 for nondominant vs −12 for dominant, p=0.017, uncorrected). There were no differences in change scores between patients with dominant or nondominant affected limbs for any of the other outcome scores.

We then performed multiple linear regression to look for predictors of either absolute scores at 6 months after treatment (T4) or the change in scores from admission to 6 months after treatment (T1−T4) using age, BI, mRS, NFI, HADS and sensation on admission together with the matching initial score on admission (eg, FM1 if attempting to predict FM4 or change from FM1 to FM4). Models predicting T4 scores for FM-UL, ARAT, CAHAI, ArmA-A and ArmA-B were all highly significant (p<0.0001). Models predicting change from T1 to T4 for ARAT, CAHAI, ArmA-A and ArmA-B (but not FM-UL) were also highly significant (p<0.0001). However, the only factor that...
Cerebrovascular disease

Figure 2  Self-reported outcome scores for all patients. Lower scores are better. Each data point represents a single patient. Top row shows individual scores at admission, discharge, 6 weeks and 6 months after discharge. Bottom row shows the individual difference scores for admission to discharge, admission to 6 weeks postdischarge and admission to 6 months postdischarge. Scores are shown for the ArmA-A and ArmA-B. Median (solid line) and upper and lower quartiles (dotted lines) are shown. ArmA-A, Arm Activity Measure A; ArmA-B, Arm Activity Measure B.

significantly contributed to these models was the matched initial score on admission (eg, ARAT at T1 predicting ARAT at T4), but not age, BI, mRS, NFI, HADS or sensation on admission. As such, this approach did not contribute further to the correlation results shown in tables 4 and 5, with regression models not able to account for a higher proportion of variance in scores at T4 or change in scores from T1 to T4 than the admission score (see also figure 3).

Relationship between outcome measures

Each outcome score assesses a different aspect of recovery. Using Spearman’s rank correlation, we examined the relationship between the changes in outcome scores from admission to 6-month follow-up. A Bonferroni correction was applied to the significance level for the 10 comparisons made between the five different outcome scores (p<0.05/10=0.005) (table 6 and figure 4). Changes in FM-UL, ARAT and CAHAI correlated weakly with one another. The ArmA-B change correlated with changes in both ArmA-A and ARAT.

DISCUSSION

We report the results from a single-centre clinical service delivering high dose and intensity of upper limb focused neurorehabilitation for people with stroke. Almost all of these patients were in the chronic (>6 months after stroke) phase. The key messages

Table 3  Changes in upper limb scores

<table>
<thead>
<tr>
<th></th>
<th>Adm - Dis</th>
<th>Adm - 6 weeks</th>
<th>Adm - 6 months</th>
</tr>
</thead>
<tbody>
<tr>
<td>FM-UL</td>
<td>6 (3 to 9)</td>
<td>8 (4 to 11)</td>
<td>9 (4 to 12)</td>
</tr>
<tr>
<td>ARAT</td>
<td>6 (2 to 8)</td>
<td>7 (2 to 10)</td>
<td>8 (3 to 11)</td>
</tr>
<tr>
<td>CAHAI</td>
<td>9 (5 to 13.5)</td>
<td>10 (5 to 16)</td>
<td>12 (6 to 17)</td>
</tr>
<tr>
<td>ArmA-A</td>
<td>−2 (−5 to −1)</td>
<td>−3 (−6 to −1)</td>
<td>−4 (−8 to −1)</td>
</tr>
<tr>
<td>ArmA-B</td>
<td>−7 (−14 to −3)</td>
<td>−10 (−17 to −4)</td>
<td>−11 (−19 to −5)</td>
</tr>
</tbody>
</table>

All scores given as median (IQR).

ARAT, Action Research Arm Test; ArmA-A, Arm Activity Measure A; ArmA-B, Arm Activity Measure B; CAHAI, Chedoke Arm and Hand Activity Inventory; FM-UL, Fugl-Meyer (upper limb).

Table 4  Correlation between baseline scores and 6-month outcomes

<table>
<thead>
<tr>
<th></th>
<th>FM-UL (T4)</th>
<th>ARAT (T4)</th>
<th>CAHAI (T4)</th>
<th>ArmA-A (T4)</th>
<th>ArmA-B (T4)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Barthel Index</td>
<td>n.s.</td>
<td>n.s.</td>
<td>0.28</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>modified Rankin Scale</td>
<td>−0.29</td>
<td>−0.25</td>
<td>−0.32</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Neurological Fatigue Index</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Hospital Anxiety and Depression Scale</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Sensation</td>
<td>n.s.</td>
<td>n.s.</td>
<td>0.22</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>FM-UL (T1)</td>
<td>0.92</td>
<td>0.84</td>
<td>0.83</td>
<td>−0.42</td>
<td>−0.63</td>
</tr>
<tr>
<td>ARAT (T1)</td>
<td>0.87</td>
<td>0.95</td>
<td>0.91</td>
<td>−0.48</td>
<td>−0.71</td>
</tr>
<tr>
<td>CAHAI (T1)</td>
<td>0.84</td>
<td>0.88</td>
<td>0.92</td>
<td>−0.46</td>
<td>−0.71</td>
</tr>
<tr>
<td>ArmA-A (T1)</td>
<td>−0.39</td>
<td>−0.44</td>
<td>−0.44</td>
<td>0.56</td>
<td>0.41</td>
</tr>
<tr>
<td>ArmA-B (T1)</td>
<td>−0.53</td>
<td>−0.58</td>
<td>−0.61</td>
<td>0.40</td>
<td>0.71</td>
</tr>
</tbody>
</table>

Correlation coefficient (rho) given only when significant at \( p<0.05 \) corrected for 55 multiple comparisons (\( p<0.0009 \)). n.s.=not significant.

ARAT, Action Research Arm Test; ArmA-A, Arm Activity Measure A; ArmA-B, Arm Activity Measure B; CAHAI, Chedoke Arm and Hand Activity Inventory; FM-UL, Fugl-Meyer (upper limb).
Figure 3  Admission score plotted against final scores at 6 months postdischarge for FM-UL, ARAT, CAHAI, ArmA-A and ArmA-B. The black line represents the line of ‘no change’. Patients above this line for FM-UL, ARAT and CAHAI have improved. Patients below this line for ArmA-A and ArmA-B have improved. FM-UL, Fugl-Meyer (upper limb); ARAT, Action Research Arm Test; CAHAI, Chedoke Arm and Hand Activity Inventory; ArmA-A, Arm Activity Measure A; ArmA-B, Arm Activity Measure B.

from our results are as follows: (i) patients were able to complete the 90 hours of the programme, despite exhibiting a range of impairments and fatigue levels, (ii) large clinical improvements in upper limb impairment (FM-UL) and activity (ARAT, CAHAI and ArmA) were observed and (iii) these changes were maintained, or even improved on, 6 months after treatment.

The QSUL programme is based on the hypothesis that a high dose and intensity of upper limb neurorehabilitation can lead to large clinically meaningful improvements in chronic stroke patients. It is worth comparing the magnitude of the changes with previous reported clinical trials of upper limb rehabilitation. Most obviously, the study by McCabe et al.10 investigated the effects of 300 hours of upper limb rehabilitation in 48 chronic stroke patients with similar characteristics to those reported here (baseline FM-UL 24, compared with 26 in our patients) and reported an increase of 8–11 points on the FM-UL.

Table 5  Correlation between baseline scores and change scores from admission to 6 months

<table>
<thead>
<tr>
<th></th>
<th>FM-UL (T1–T4)</th>
<th>ARAT (T1–T4)</th>
<th>CAHAI (T1–T4)</th>
<th>ArmA-A (T1–T4)</th>
<th>ArmA-B (T1–T4)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Barthel Index</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>modified Rankin Scale</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Neurological Fatigue Index</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Hospital Anxiety and Depression Scale</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>Sensation</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>FM-UL (T1)</td>
<td>n.s.</td>
<td>0.40</td>
<td>0.45</td>
<td>n.s.</td>
<td>−0.24</td>
</tr>
<tr>
<td>ARAT (T1)</td>
<td>n.s.</td>
<td>0.36</td>
<td>0.52</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>CAHAI (T1)</td>
<td>n.s.</td>
<td>0.42</td>
<td>0.35</td>
<td>n.s.</td>
<td>n.s.</td>
</tr>
<tr>
<td>ArmA-A (T1)</td>
<td>n.s.</td>
<td>n.s.</td>
<td>−0.31</td>
<td>−0.72</td>
<td>n.s.</td>
</tr>
<tr>
<td>ArmA-B (T1)</td>
<td>n.s.</td>
<td>−0.26</td>
<td>−0.38</td>
<td>−0.28</td>
<td>−0.26</td>
</tr>
</tbody>
</table>

Correlation coefficient (rho) given only when significant at p<0.05 corrected for 55 multiple comparisons (p=0.0009). n.s.=not significant.

ARAT, Action Research Arm Test; ArmA-A, Arm Activity Measure A; ArmA-B, Arm Activity Measure B; CAHAI, Chedoke Arm and Hand Activity Inventory; FM-UL, Fugl-Meyer (upper limb).
at the end of 300 hours of upper limb treatment (although later follow-up results have not been reported). In the QSUL programme, the median FM-UL (modified) improvement was 6 points after the 90 hours programme, increasing to 9 points by 6-months follow-up, achieving a similar overall magnitude of change in comparison to McCabe et al. In addition to reductions in impairment (FM-UL), our patients also improved at the level of activity (ARAT and CAHAI) and participation (using the patient-reported outcome measure, ArmA). Importantly, we report that these group-level effects continued to improve at 6 weeks and then 6 months after treatment had stopped, (figures 1 and 2).

Other than the study by McCabe et al., the general view has been that large changes, especially in impairment, have not been possible to deliver such high doses in current healthcare settings (the pragmatic view), it is in fact precisely the role of clinical research to challenge what we currently do in order to reshape our clinical services to make them better (the aspirational view). Those involved in stroke medicine will recognise that this is exactly the effect that thrombolysis had on acute management of stroke, leading to the high-quality hyperacute stroke services we see today. Post-stroke neurorehabilitation must adopt the same aspirational approach.

While the appropriate dose of upper limb rehabilitation is debated, the content of the intervention receives less attention. Our approach was based on analysis of movement and performance in activities of daily living, reduction of impairment and re-education of quality and control of movement within activities of daily living, all performed in an iterative and nonlinear fashion. Education, self-efficacy and goal setting were integral components of the programme, and we can speculate that the education and focus on self-efficacy contributed to the continued increase in scores after treatment stopped. The therapeutic approach taken towards specific impairments was the same across patients, but approaches at the activity and participation level will necessarily vary as they are tailored to an individual’s specific meaningful goals. We acknowledge that an immediate challenge for us and for the field in general is to be able to define what the ‘active ingredients’ of neurorehabilitation are, so that these elements can be further tested. Our approach appears to have similarities with that taken by McCabe et al., with an emphasis on the quality of the movement as well as the final goal of the movement. The latter is often the sole focus of task-specific repetition, which has been used in many upper limb rehabilitation studies to date. The evidence for repetitive task-specific training is of low to moderate quality and the effects have been disappointingly small even when the number of repetitions has been greatly increased. It is difficult to tell whether this lack of effect is specific to task-specific training because in general the dose (in terms of hours of active training) has been quite low, whether delivered by a therapist or robotic device. There have been suggestions that chronic stroke patients might not tolerate more training than has been offered in these trials, but the study of McCabe et al. and now our own clinical experience undermine this view.

Although this result is exciting, we must acknowledge that it is a single-centre clinical service with no randomisation and no control group. Although there are some reports of late improvement up to 12 months after lacunar stroke, we expect that chronic stroke patients, who are often several years after their stroke, will not change across this range of outcome scores without treatment. We suggest that it is unlikely that this magnitude of change would be seen without therapeutic intervention, but this could be possible if the dose (measured in time) of the intervention may have been much lower than we report here. While this report of our single-centre study was based on analysis of movement and performance in activities of daily living, reduction of impairment and re-education of quality and control of movement within activities of daily living, all performed in an iterative and nonlinear fashion. Education, self-efficacy and goal setting are integral components of the programme, and we can speculate that education and focus on self-efficacy contributed to the continued increase in scores after treatment stopped. The therapeutic approach taken towards specific impairments was the same across patients, but approaches at the activity and participation level will necessarily vary as they are tailored to an individual’s specific meaningful goals. We acknowledge that an immediate challenge for us and for the field in general is to be able to define what the ‘active ingredients’ of neurorehabilitation are, so that these elements can be further tested. Our approach appears to have similarities with that taken by McCabe et al., with an emphasis on the quality of the movement as well as the final goal of the movement. The latter is often the sole focus of task-specific repetition, which has been used in many upper limb rehabilitation studies to date. The evidence for repetitive task-specific training is of low to moderate quality and the effects have been disappointingly small even when the number of repetitions has been greatly increased. It is difficult to tell whether this lack of effect is specific to task-specific training because in general the dose (in terms of hours of active training) has been quite low, whether delivered by a therapist or robotic device. There have been suggestions that chronic stroke patients might not tolerate more training than has been offered in these trials, but the study of McCabe et al. and now our own clinical experience undermine this view.

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assertion requires empirical confirmation. Our work, however, provides strong justification for undertaking such high-dose clinical trials. Clinic data on which to base these future studies are sorely lacking and this has led to the proliferation of low-dose pragmatic rehabilitation studies. However, there is still much to be learnt from published data arising from clinical services in neurorehabilitation, making up for lack of clinical trial rigour with large sample sizes that are rarely achieved in clinical trials. Data from clinical services therefore have the potential to refine what experimental questions are addressed in future randomised clinical trials. Another limitation is the unblinded nature of the assessments, introducing possible bias into the results. All assessors are also therapists on the programme, although we ensure that assessments are carried out by therapists from another team who are not treating the patient. Fidelity of outcome scores is always an important issue in clinical trials. In our programme, we have instigated regular in-service training for FM-UL, ARAT and CAHAI to ensure high levels of reproducibility between assessors.

A common complaint regarding single-centre clinical services is that they select only patients most likely to succeed. It is important to acknowledge that the QSUL programme does not have strict inclusion/exclusion criteria. Although our patients are generally younger than the average stroke patient and have high levels of independence, they have a wide range of upper limb impairments. It can be seen from figures 1 and 2 that the range of severity of the upper limb on the QSUL programme was very broad, quite unlike a clinical trial, although the median scores suggest that patients were weighted towards the moderate–severe end of upper limb impairment and activity limitation. We would argue that the QSUL programme is aspirational in its goals (high-quality, high-dose, high-intensity treatment) and pragmatic in its application (wide range of patients admitted onto the programme). It is interesting to consider the potential impact of initial upper limb severity on our outcomes, but in fact, although initial score correlates well with the final score at 6 months (table 4), the change in score correlates only weakly with initial score (table 5, figure 3). There are two ways this ‘mixed-bag’ approach may have potentially reduced the overall magnitude of the clinical effects we observed. First, there may be a ceiling effect from patients with relatively high scores on admission. Second, many of our patients were quite severely impaired. Indeed, a quarter of our patients had an admission ARAT below 7, which suggests they have great difficulty in using their hand in any kind of activity of daily living. The treatment goals in these type of patients might initially be very different to the moderate-mild type patients, with benefits less likely to be reflected in changes in FM-UL and ARAT compared with the CAHAI or ArMA, for example. However, the wide range of patients treated in the programme means that no clear patterns, in relation to differential effects on various scores, could be seen (table 6, figure 4).

Some might argue that the wide range of responses, including those patients who changed very little, points to the need for stratification and selection of those patients most likely to respond, and this is certainly a reasonable view. However, the first point to make is that our results show that with large enough numbers and a high enough dose of the intervention, it is possible to demonstrate large group-level effects in relatively unslected populations of chronic stroke patients. Indeed, if the former (stratification) approach is to be adopted, then these kinds of large data sets will be required to determine what the characteristics of the ‘responders’ are likely to be. The second point to make is that we found no strong clinical indicators, among our measured explanatory variables, of which patients were most likely to change, including sensory loss in the hand, fatigue and depression (tables 4 and 5). It might be that neurophysiological (presence or absence of motor evoked potentials), neuroimaging (assessment of corticospinal tract, whole brain damage or brain function) or cognitive (sustained attention, memory, motivation) measures are required for accurate stratification based on likely response, and this can certainly be tested in future.

In summary, here we present the results from a single-centre clinical service dedicated to post-stroke upper limb neurorehabilitation in chronic stroke patients. Despite treating patients with a wide range of impairment, we were able to show a large group-level magnitude of change at both the impairment and activity level. Our experience suggests that much higher doses and intensity of upper limb neurorehabilitation can be delivered and our results should inform future clinical trial design.

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