Effect of clinical guidelines in nursing, midwifery, and the therapies: a systematic review of evaluations

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Abstract

Background—Although nursing, midwifery, and professions allied to medicine are increasingly using clinical guidelines to reduce inappropriate variations in practice and ensure higher quality care, there have been no rigorous overviews of their effectiveness. 18 evaluations of guidelines were identified that meet Cochrane criteria for scientific rigor.

Methods—Guideline evaluations conducted since 1975 which used a randomised controlled trial, controlled before and after, or interrupted time series design were identified through a combination of database and hand searching.

Results—18 studies met the inclusion criteria. Three studies evaluated guideline dissemination or implementation strategies, nine compared use of a guideline with a no guideline state; six studies examined skill substitution: performance of nurses operating according to a guideline were compared with standard care, generally provided by a physician. Significant changes in the process of care were found in six out of eight studies measuring process and in which guidelines were expected to have a positive impact on performance. In seven of the nine studies measuring outcomes of care, significant differences in favour of the intervention group were found. Skill substitution studies generally supported the hypothesis of no difference between protocol driven by nurses and care by a physician. Only one study included a formal economic evaluation, with equivocal findings.

Conclusions—Findings from the review provide some evidence that care driven by a guideline can be effective in changing the process and outcome of care. However, many studies fell short of the criteria of the Cochrane Effective Practice and Organisation of Care Group (EPOC; formerly the Cochrane Collaboration on Effective Professional Practice).

Keywords: clinical guidelines; nursing; midwifery; systematic review; clinical effectiveness
Methods

INCLUSION AND EXCLUSION CRITERIA
The definition of clinical guidelines by Field and Lohr[1] as “systematically developed statements to assist practitioner decisions about appropriate health care for specific clinical circumstances” was used. We were interested in all types of guideline, not simply those based on evidence.

To be eligible for inclusion in our review, a study had to be an evaluation of the effect of a clinical guideline on the behaviour of nurses or other professionals allied to medicine or on patient outcomes. We included studies which evaluated an entire, or identified component, of a guideline aimed at these professions and which specified which part or parts of the guideline should be implemented by the profession in question. We considered not only studies of guidelines developed by the target profession for their own use, but also those in which the guideline was developed or disseminated by another professional group but targeted the behaviour of nurses or other professionals allied to medicine.

We included both studies which evaluated the effectiveness of guidelines against a non-intervention control (routine care) and those comparing different dissemination and implementation strategies of a guideline—for example, dissemination strategy X versus dissemination strategy Y. Studies which simply described the development, dissemination, or implementation of a guideline, without an evaluation of the impact, were excluded, as were studies in which <50% of recipients of the guideline were nurses, midwives, or other professions allied to medicine and studies in which it was not possible to separate the effects of guidelines on the behaviour of professionals from the impact on the behaviour of other health professionals. A start date of 1975 was imposed, firstly to make our review comparable with that of Grimshaw and Russell,[2,3] and secondly because MEDLINE does not contain abstracts before that year.

METHODOLOGICAL CRITERIA
To be compatible with Grimshaw et al.[2,3] and in line with standard practice for EPOC reviews, three types of study designs were included.

(1) Randomised controlled trials, where we distinguished between simple randomised controlled trials (in which the unit of allocation is the patient) and designs in which the unit of allocation is the health professional and which are therefore less susceptible to contamination.

(2) Controlled before and after studies.

(3) Interrupted time series designs, if there was a clearly defined point in time when the intervention occurred and at least two data collection points before and two after the intervention.

SYSTEMATIC REVIEW OF LITERATURE
We searched: (a) computerised databases (MEDLINE; CINAHL (Cumulative Index of Nursing and Allied Health Literature); PsycLIT; EMBASE; NEED (NHS Economic Evaluations Database); DARE (Database of Abstracts of Reviews of Effectiveness); DHSS-DATA; SIGLE; and the National Research Register); (b) the EPOC register of studies; (c) citations in papers identified, and (d) Quality in Health Care*. We also contacted content area experts, the libraries of appropriate professional bodies—for example, the Royal College of Nursing—and various mailbases on the internet—for example, evidence-based-health@mailbase.ac.uk.

Search strategies for electronic databases were developed sequentially, starting with MEDLINE, as this was expected to yield the highest number of relevant papers. We adapted the focus section of the EPOC search strategy (designed to identify randomised trials, controlled before and after and interrupted time series studies) to expand the section on clinical guidelines. We searched for guidelines and various synonyms in titles and abstracts as well as carrying out keyword searches. MEDLINE also identifies “publication types”, which allowed us to search for papers tagged as “practice guideline” or “guideline”. We then worked with an information scientist to adapt the search strategy for use with the other databases.

SCREENING OF IDENTIFIED PAPERS
The full text version of each study identified as potentially relevant to our review was read by one reviewer (NC, EM, NR, JS, or LHT) and assessed against several criteria with a decision tree developed for this review. To establish consistency between raters, all reviewers initially completed checklists for the same five papers; discrepancies were discussed and resolved. All those studies identified by one reviewer as meeting all the inclusion criteria, and those in which there was an element of doubt, were referred to a second reviewer; discussion between the two reviewers took place to resolve any remaining differences.

DATA EXTRACTION
We used the checklist and data extraction template developed by EPOC to abstract data from studies meeting our inclusion criteria. Data extraction was undertaken by two reviewers working independently and included both methodological details—for example, units of allocation and data analysis; methodological quality criteria—and information about study participants, interventions, outcomes, and results. The two reviewers then met to compare their judgements and any discrepancies were discussed and resolved. A “master version” of the data extraction template was agreed and the details were put into an Idealist[4] database.

* Quality in Health Care is now included in the following databases: HEALTH (Health Planning and Administration) Database of the National Library of Medicine and American Hospital Association Resource Center from 1995 onwards, CINAHL, ISI Science Citation Index (public health, environmental and occupational), ISI Current Contents (clinical medicine), and EMBASE.
DATA ANALYSIS
As the heterogeneity of clinical areas, design of studies, source and format of interventions, outcomes measured, and participating health professionals, was substantial, an overall estimate of effect was likely to have little practical meaning. We therefore opted simply to report the effects on the process and outcome of care in the same way as they were reported in the original papers, rather than to attempt comparisons with some common unit of change.

Results
We identified 18 studies which met our inclusion criteria. In all but one, guidelines were aimed, in part at least, at nurses. Of these studies, in five cases, physicians were also targeted. The study by Franz et al was aimed solely at dietitians. Shaffer and Wexler presented a truly multi-disciplinary guideline, aimed at an entire lipid clinic team comprising nurses, pharmacists, psychologists, and dietitians.

METHODOLOGICAL QUALITY
We identified 13 randomised controlled trials. In three studies the unit of allocation was the health professional or provider unit. In all three, the randomisation process was not described and it is not clear whether there was blinded assessment of outcomes. One trial had a unit of analysis error: the unit of randomisation was the provider, but the unit of analysis was the patient. Such unit of analysis errors artificially increase the power of the study and may lead to misleadingly significant findings.

There were 10 randomised controlled trials in which the unit of allocation was the patient. Six of these examined the effect of substituting another health professional (usually a nurse) working to protocols for usual care provided by a doctor: obviously in these studies, randomisation by health professional would not be possible. There were major differences in methodological quality: all 13 studies had good follow up; however, just over half (7/13) blindly assessed outcomes and reliably measured at least some key outcomes (7/13). A few studies reported method of randomisation.

We identified two controlled before and after studies in their evaluation of opinion leaders and implementation strategies. All were randomised controlled trials. Within these studies, the interventions were heterogeneous, precluding combining the findings from two or more studies. Seto et al compared three groups: all received guidelines; one received both lectures and input from local opinion leaders; the remaining two received either lectures or input from opinion leaders. Some evidence was found that the combined strategy of opinion leaders and lectures led to higher compliance with correct practice than did either strategy alone. The weakest strategy was a lecture alone.

Three studies were interrupted time series designs. In the study by French et al, the intervention seemed to be independent of other changes during the study period; in the study of Mitchell and Jones no details were provided as to whether this was or was not the case. In the study of Tilden and Shepherd, in which the targeted behaviour was identification of battered women, the authors stated that media coverage and an initiative by hospital administration (aimed at all staff) to increase the level of detail in patient documentation may have intensified the effect of the intervention. No interrupted time series study had ≥ 12 data points before and after the intervention; only one study did a formal test for trend. All interrupted time series studies were prone to detection bias. In two studies data collection methods were identical before and after the intervention. However, in the study of Mitchell and Jones data on rates of falls before intervention were collected from retrospective analysis of incident forms of falls, and prospective data were collected by adding a “falls reporting document” to the incident form. As this seemed to be part of the intervention, it directly influenced data collection. Whether the intervention directly influenced data collection in the two other interrupted time series studies is not clear. Outcomes were not assessed blindly in one study; in the remaining two studies it is not clear whether this was the case. In only one study was information provided about the completeness of the data set: all adult female trauma patients were included. No interrupted time series study measured outcomes reliably.

STUDIES EVALUATING DISSEMINATION AND IMPLEMENTATION STRATEGIES
Three studies (table) explicitly evaluated guideline dissemination or implementation strategies. All were randomised controlled trials. Within these studies, the interventions were heterogeneous, precluding combining the findings from two or more studies. Seto et al in their evaluation of opinion leaders and in-service lectures as methods of disseminating a guideline on urinary catheter care to nurses, compared three groups: all received guidelines; one received both lectures and input from local opinion leaders; the remaining two received either lectures or input from opinion leaders. Some evidence was found that the combined strategy of opinion leaders and lectures led to higher compliance with correct practice than did either strategy alone. The weakest strategy was a lecture alone.

Herman et al found that a “nurse protocol” intervention (primarily an implementation strategy) led to positive changes in the process of care (offer of influenza and pneumococcal vaccine). There were three comparison groups: all received guidelines, educational materials,
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<th>Study and implementation strategies:</th>
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<tr>
<td>Herman et al (1994)** RCT-Pr US</td>
<td>Nurses (number not specified) and 66 doctors in 3 ambulatory medical practices Provider: 1 practice in each of 3 arms Proportion of patients followed up: 100% of eligible patients for influenza vaccination group; 75% for pneumococcal vaccination group</td>
<td>Pneumococcal and influenza vaccinations</td>
<td>I: Guidelines; educational materials; lectures I2: As 1; provider and patient education provided at each clinic visit I3: As 1; plus protocol</td>
<td>Offer of influenza vaccination: I1: 48% (129/271) I2: 50% (122/242) I3: 68% (166/243) p&lt;0.006 Offer of pneumococcal vaccination: I1: 5.4% (18/369) I2: 6.5% (19/292) I3: 28.3% (89/314) p&lt;0.001</td>
<td>Influenza vaccination: I1: 5.9%; I2: 5.8%; I3 13.2% p&lt;0.003 Pneumococcal vaccination I1: 2.0%; I2: 1.4%; I3: 6.7% p&lt;0.001</td>
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<td>McDonald et al (1980)** RCT-Pr (Crossover trial) US</td>
<td>Nurses and 26 doctors in 1 medical outpatient clinic 3-Treatment crossover design: all providers experienced all conditions in 1 of 6 possible different treatment sequences Proportion of patients followed up: not applicable</td>
<td>Use and follow up of medications</td>
<td>I: Computer reminders I2: Computer reminders + bibliographic citations C: Standard practice</td>
<td>Individual provider response rate = number of appropriate responses/total number possible: average nurse response rate to prompts I1: 30%; C: 25% (n=5 nurses) Not significant: insufficient data provided to calculate p values or 95% CIs</td>
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<td>Seeto et al (1991)** RCT-Pr Hong Kong</td>
<td>220 Nurses in 1 hospital Provider: 1 ward in each of 3 arms Proportion of patients followed up: not applicable</td>
<td>Urinary catheter care</td>
<td>I: Guidelines; local opinion leaders; lecture I2: Guidelines; local opinion leaders C: Guidelines; lecture</td>
<td>Change in reported practice, 2 weeks before v 3 weeks after intervention: I1: 5.63 (50%) I2: 4.96 (35%) C: 5.29 (38%) Mean change in C significantly lower than I1 and I2 p&lt;0.05 I1 v I2 not significant: insufficient data provided to calculate p values or 95% CIs Percentage of correct catheter practices, 1 month after intervention: I1 v I2 p&lt;0.05 I1 v C: p&lt;0.05</td>
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Comparison of guideline v no guideline: Franz 1995** RCT-Pr US

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<td>Study and implementation strategies:</td>
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<td>Non-insulin dependent diabetes mellitus</td>
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Comparison of guideline v no guideline: Frigoletto et al (1994)** RCT-Pr US

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<td>Two maternity units in 1 hospital Patients: 1017 intervention 917 control Protocol eligible sub-group: 678 intervention 585 control Proportion of patients followed up: 99% in total</td>
<td>Delivery of first babies</td>
<td>I: Protocol (midwives) C: Standard care</td>
<td>Frequency of caesarean section: I1: 19.5% (197/1009); I2: 19.4% (176/906) RR = 1.0 95% CI (0.8–1.2) Mean frequency of vaginal examination (hours†): I1: 1.6; C: 2.3 Artificial rupture of membranes: I1: 61%; C: 51% RR = 1.259 CI (1.1–1.3) Rupture within 1 hour of admission: I1: 76%; C: 15% RR = 17.8 95% CI (13.3–23.8) Administration of oxytocin: I1: 70%; C: 56% RR = 1.395 CI (1.2–1.4) Administration of epidural anaesthesia: I1: 54%; C: 64% RR = 0.8 95% CI (0.8–0.9) Rate of caesarean section: I1: 19.5% (197/1009); I2: 19.4% (176/906) for all patients RR = 1.0 95% CI (0.8–1.2) I: 10.9%; C: 11.5% for protocol eligible group RR = 0.9 95% CI (0.4–1.9) Duration of labour (protocol eligible subgroup): I: 6.2; C: 8.9 hours Insufficient data provided to calculate 95% CIs Rate of maternal fever (protocol eligible subgroup): I: 7%; C: 11% p&lt;0.007 Rate of other maternal outcomes: no significant differences in foetal distress, placental abruptio, shoulder dystocia, vaginal lacerations Duration of labour &gt;12 hours (protocol eligible subgroup): I: 9%; C: 26% p&lt;0.0001 Infant outcomes: no significant differences in frequency of jaundice, seizures, treatment for sepsis, resuscitation at birth or admission to neonatal ICU</td>
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*Clinical area: Delivery of first babies, Delivery, Delivery of babies, Urinary catheter care, Delivery of babies, Delivery of first babies, Delivery of first babies, Delivery of babies, Delivery of first babies, Delivery of babies, Delivery of babies, Delivery of first babies, Delivery of first babies.
### Comparison tables (continued)

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<tr>
<td><strong>Naylor</strong> 1990&lt;sup&gt;10&lt;/sup&gt; RCT/Pt US</td>
<td>2 Nurses (in intervention group) in 1 hospital</td>
<td>Patients: 20 intervention 20 control</td>
<td>Discharge planning for &gt;70 year-olds in medical and surgical units</td>
<td>I: Protocol C: Standard practice</td>
<td>Mean duration of stay (days): I: 8.2 ± 4.9; C: 9.05 ± 7.66; not significant: insufficient data to calculate p value or 95% CIs. Post discharge infection rates: I: 33.3% (6/20); C: 50% (10/20); RR = 0.43 (95% CI 0.11–1.57). Post discharge rates of readmissions to hospital: I: 16.7% (4 readmissions, 3 patients); C: 64.7% (12 readmissions to hospital, 11 patients); p = 0.05. Results reported for medical and surgical diagnostic related group Medical and surgical separately. Mean duration of initial admission to hospital (days): Medical: I: 7.4 (3.8); C: 7.5 (5.2) – difference (95% CI) 0.1 (1.6 to 1.4). Surgical: I: 15.8 (9.4); C: 14.8 (8.3) – difference (95% CI) 1.0 (–2.0 to 4.0). Mean days to readmission to hospital: I: 45.6; C: 31; p = 0.12. Rate of readmission to hospital: Medical: I: (15%); C: (16%); 11 difference (95% CI) –1% (–8% to 12%). Surgical: I: (10%); C: (7%); 5 difference (95% CI) –3% (–7% to 13%). Mean duration of readmission to hospital: Medical: I: 94; C: 100; difference (95% CI) –6 (–83 to 71). Surgical: I: 52; C: 26 difference (95% CI) 26 (–8 to 60). Occurrence of iv related phlebitis: I: increase from 33.5% to 20.9% and p = 0.05; C: decrease from 23.8% to 26.7%; p = 0.25. Frequency of bacterial colonisation of infusion devices: I: increase from 12.7% to 19.4% and p = 0.25; C: decrease from 7.1% to 5.9% and p = 0.90. Patient comfort with IV insertion: not reported in appropriate format.</td>
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<td><strong>Naylor</strong> 1994&lt;sup&gt;10&lt;/sup&gt; RCT/Pt US</td>
<td>2 Nurses and doctors (number not specified) in 1 hospital</td>
<td>Patients: 364 patients randomised; 276 analysed; 140 intervention 136 control</td>
<td>Discharge planning for &gt;70 year-olds in medical and surgical units</td>
<td>I: Protocol C: Standard practice</td>
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<td><strong>Larson and Hargis</strong> 1984&lt;sup&gt;6&lt;/sup&gt; CBA US</td>
<td>10 nurses (in intervention group) and doctors (number not specified) in 2 intervention and 3 control wards in 1 university affiliated hospital</td>
<td>Patients: 707 (in total) Proportion of patients followed up: not clear</td>
<td>Intravenous therapy</td>
<td>I: Guidelines; outreach visits C: Standard practice</td>
<td>Percentage of items documented correctly: I: increase from 57% to 79%; C: increase from 52.5% to 68.5%; p &lt; 0.001 Test inappropriate</td>
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<td><strong>Shaffer and Weeler</strong> 1995&lt;sup&gt;5&lt;/sup&gt; CBA US</td>
<td>1 lipid clinic team (nurse led) and 1 general medicine clinic team in 1 medical centre Provider: 1 intervention 1 control Patients: 60 intervention 60 control</td>
<td>I: Guidelines (nurse, clinical pharmacist, dietitian, clinical psychologist) C: No intervention (doctors)</td>
<td>Hyperlipidaemia</td>
<td>Percentage of patients for whom a new drug was prescribed: I: 85% (51/60); C: 18% (11/60); p &lt; 0.001 Number of patients in whom an existing drug was stopped: I: 22 patients; C: 3 patients. p &lt; 0.001 Percentage of patients receiving expert dietary counselling: I: 100% (60/60); C: 42% (25/60) p &lt; 0.001</td>
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<td>French et al 1989&lt;sup&gt;9&lt;/sup&gt; IIS Hong Kong</td>
<td>3 Infection control nurses in 1 hospital</td>
<td>Patients: ranged from 809 to 1260 in 7 prevalence surveys</td>
<td>Infection rates</td>
<td>I: Guidelines C: No intervention</td>
<td>Total cholesterol at follow up: I: 6.08 mmol/l (from 7.65); C: 6.72 mmol/l (from 7.50; p = 0.004. LDL-cholesterol at follow up: I: 3.78 mmol/l (from 5.20); C: 4.60 mmol/l (from 5.17); p &lt; 0.001. HDL-cholesterol at follow up: I: –1.03 mmol/l (from 1.11); C: –1.14 mmol/l (from 1.19); not significant: insufficient data to calculate p value or 95% CIs. Triglycerides at follow up: I: –3.39 mmol/l (from 4.13); C: –3.56 mmol/l (from 4.45); not significant: insufficient data to calculate p value or 95% CIs. Body mass index: I: 28.3 kg/m² (from 27.9 kg/m²); C: 28.5 kg/m² (from 28.8 kg/m²); not significant: insufficient data to calculate p value or 95% CIs. Percentage achieving national cholesterol educational programme LDL-cholesterol goal (3.36 mmol/l): I: 44% (19/43); C: 11% (3/28); RR = 4.1 95% CI (1.4–12.7). Percentage on lipid lowering drug: I: 69% (51/75); C: 35% (21/60); RR = 3.75 95% CI (2.05–6.87). Pre intervention hospital acquired infection rates (crude) 8.9%, 10.5%. Pre intervention hospital acquired infection rates (crude) 7%, 7.8%, 6.2%, 6%, 5.6% p &lt; 0.001. Pre intervention hospital acquired infection rates (adjusted) 8.5%, 9.9%. Pre intervention hospital acquired infection rates (adjusted) 6.7%, 7.5%, 6.7%, 6.0%, 6.2% p &lt; 0.001. Hospital acquired urinary tract infection: I: 3.1%; C: 3.3%. p = 0.13. Pre intervention hospital acquired infection rates (crude) 8.9%, 8.5%. Pre intervention hospital acquired infection rates (adjusted) 6.7%, 7.5%, 6.7%, 6.0%, 6.2% p &lt; 0.001. Pre intervention hospital acquired infection rates (crude) 8.9%, 8.5%. Pre intervention hospital acquired infection rates (adjusted) 6.7%, 7.5%, 6.7%, 6.0%, 6.2% p &lt; 0.001. Hospital acquired urinary tract infection: I: 3.1%; C: 3.3%. p = 0.13.</td>
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<td><strong>Mitchell and Jones</strong> 1996&lt;sup&gt;7&lt;/sup&gt; IIS Australia</td>
<td>Nurses (number not specified) on 1 medical ward in an acute care hospital</td>
<td>Falls I: Protocol C: No intervention</td>
<td>Falls</td>
<td>Increase in rate of identification of positive battering from 7 cases (9.72% of total) to 17 cases (22.97% of total) p = 0.03 OR likelihood of identification 2.74 times greater post-intervention</td>
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<td><strong>Tilden</strong> 1987&lt;sup&gt;4&lt;/sup&gt; IIS US</td>
<td>22 Nurses in 1 hospital emergency department</td>
<td>Female adult trauma</td>
<td>I: Guidelines; staff education programme; local consensus processes C: No intervention</td>
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<td><strong>Skill-substitution:</strong></td>
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<td>Greenfield et al 1975&lt;sup&gt;23&lt;/sup&gt; RCT-Pt (Cross over trial) US</td>
<td>1 nurse and 13 doctors in 1 clinic. Patients: 151 randomised; 146 intervention; 70 control</td>
<td>Management of dysuria, frequency, and vaginal discharge</td>
<td>I: Protocol (nurses); C: No intervention (doctors)</td>
<td>Concordance between nurse and physician on medical history: 139/146 (95.2%) CI 95% (91.8–98.7%). Concordance on results of physical examination: 137/146 (93.8%) CI 95% (89.8–97.5%). Concordance on therapy and referral: 144/146 (99%) All referrals appropriate; protocol did not fail to recommend referral where there should have been one Laboratory agreed with nurse in (93%) 54/58 urinalyses 95 CI 93.1% (86.5–99.6%).</td>
<td>Alleviation or improvement of symptoms: I: (57%) 66/68; C: (97%) 63/65 p&lt;1.00 Results of treatment with antibiotics (urine samples sterile): I: (71%) 10/14; C: (93%) 8/15 p=0.45</td>
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<tr>
<td>Greenfield et al 1975&lt;sup&gt;23&lt;/sup&gt; RCT-Pt US</td>
<td>5 nurses and 32 doctors in 1 walk in clinic. Patients: 592 randomised 419 completed study; 221 Intervention, 197 Control</td>
<td>Management of low back pain</td>
<td>I: Protocol (nurses); C: No intervention (doctors)</td>
<td>Number of x ray films ordered: I: 21; C: 36 p&lt;0.01 Initial or final diagnosis: no significant differences except in initial diagnosis of other (I: 10.8%; C: 1%) p&lt;0.01</td>
<td>I patients significantly more satisfied than C patients p&lt;0.005 Symptom relief: resolved or improved in I: 71 (144/203); C: 74.6% (144/193) p=0.43 Complications: no patient managed by nurses alone developed serious complications Return with back problems within 3 months: I: 27%; C: 18% p=0.04</td>
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<td>Jewell and Hopc1988&lt;sup&gt;24&lt;/sup&gt; RCT-Pt UK</td>
<td>1 Nurse and 1 doctor in 1 general practice. Patients: 17 intervention 19 control</td>
<td>Hypertension</td>
<td>I: Protocol (nurses); I2: Protocol (doctors)</td>
<td>Percentage of patients with record of: Smoking habit (once): I1: (100%) 15; I2: (84%) 16 patients p=0.32 Urine test (once): I1: (93%) 14; I2: (21%) 4 patients p&lt;0.0002 Blood pressure (at each visit): I1: (100%) 90; I2: (100%) 103 visits p=1.00 Pulse (at each visit): I1: (98%) 88; I2: (65%) 67 visits p&lt;0.01 Weight (at each visit): I1: (90%); I2: (37%) 38 visits p&lt;0.001</td>
<td>Symptom relief: resolved or improved: I: 73% (162/222); C: 68.5% (135/197) p=0.33 Patient satisfaction: satisfaction with 6 of 9 elements significantly higher in intervention group; no significant difference for remaining 3 elements; p ranged from 0.3 to 0.78. No significant differences in total satisfaction except in initial diagnosis of other (I: 10.8%; C: 1%) p&lt;0.01 Knowledge of medication: drug names recalled: I1: 11; I2: 20 p=1.00 Drug names not recalled: I1: 12; I2: 17 Patient satisfaction: number reporting satisfaction: I1: 15/17; I2: 17/19 p=1.00 All nurses protocol group chose to return to doctor care; only 3 patients in both groups did not wish to be looked after by a nurse</td>
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<td>Klassen et al 1993&lt;sup&gt;25&lt;/sup&gt; RCT-Pt Canada</td>
<td>21 Triage nurses and doctors (number not specified) in 1 emergency room. Patients: 491 intervention 494 control</td>
<td>Radiology referrals</td>
<td>I: Protocol (nurses); C: No intervention (doctors)</td>
<td>Percentage of patients having x ray films ordered: I: 81.9%; C: 87.1% p&lt;0.03 Positive findings as percentage of ordered x ray films: I: 40.8%; C: 42.6% p=0.21. Missed positive findings as percentage of whole group: I: 3.2%; C: 3% p=0.001 Mean time spent in the emergency department: I: 3.3 hours; C: 3.6 hours p=0.001</td>
<td>Mean postoperative blood loss: I: 817 ml; C: 976 ml p=0.08 Mean activated partial thromboplastin time: I: 31.7 s; C: 47.3 s p=0.01 Robustion emergency care required: I: 4% (4/108); C: 8% (12/146) p=0.32</td>
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<td>Zeller et al 1992&lt;sup&gt;26&lt;/sup&gt; RCT-Pt Australia</td>
<td>Nurses and doctors (number not specified) working in 1 cardiothoracic intensive care unit. Patients: 159 intervention 163 control</td>
<td>Management of postoperative bleeding after cardiac surgery</td>
<td>I: Guideline (nurses); C: No intervention (doctors)</td>
<td>Number of patients in whom coagulation tests performed: I: 13% (14/108); C: 14% (20/146) p=0.40 Mean time to coagulation test: I: 291 minutes; C: 454 minutes p=0.30 Volume of packed cells infused: presented graphically p&lt;0.01 Volume of total blood and blood products infused: presented graphically p&lt;0.05</td>
<td>Mean postoperative blood loss: I: 817 ml; C: 976 ml p=0.08 Mean activated partial thromboplastin time: I: 31.7 s; C: 47.3 s p=0.01 Number requiring emergency care: I: 4% (4/108); C: 8% (12/146) p=0.32</td>
</tr>
</tbody>
</table>

RCT-Pt=randomised controlled trial by provider; RCT-Pt=randomised controlled trial by patient; CBA=controlled before and after study; ITS=interrupted time series study; I=intervention; I1, I2, I3=intervention 1, intervention 2, intervention 3; C=control; LDL-cholesterol=low density lipoprotein cholesterol; HDL-cholesterol=high density lipoprotein cholesterol; RR=risk ratio; BP=blood pressure; OR=odds ratio.

and lectures; group two received additional provider and patient education at each clinic visit; and group three received an additional protocol. This was a complex intervention, comprising a flowchart record incorporating the recommendations of the guideline—that is, a patient specific prompt—and a redefinition of the nurse’s role. Herman et al<sup>27</sup> found that an
active dissemination strategy (involving provider education) had a minor effect on professional behaviour in the group receiving only guidelines, educational materials, and lectures. The impact of the guidelines on outcomes of care was also assessed: a significant difference between intervention and control groups was found, but in the opposite direction to that which was expected: the percentage of patients declining immunisation increased in the intervention period to 13.2% in the prevention team group, 5.8% in the patient education group, and 5.9% in the control group. However, there was an error in the unit of analysis in this study: hence findings were more likely to be significant.

McDonald et al.7 compared the effect of computerised prompts specific to the target profession at the time of consultation, computerised prompts plus bibliographic citations, and standard care on use and follow-up of medications. However, the results for nurses were presented with the two intervention groups combined, so it is not possible to compare the two methods of implementation. With the computerised prompts and computerised prompts plus citations groups combined, the intervention did not seem to lead to a significant change in nurses’ behaviour; this finding was contrary to that for physicians in the same study. It is possible that this negative finding was due to a lack of power to detect a difference (n=five nurses).

GUIDELINE VERSUS NO GUIDELINE STUDIES
Nine studies (table) compared the introduction of guidelines with a no guideline control.20–24 26-33 These studies aimed to improve the quality of care delivered by the target profession. Of these, four were randomised controlled trials,21 24 29 30 three were interrupted time series studies,22 23 31 and two were controlled before and after studies.22 31 Targeted behaviours were nutrition therapy for patients with non-insulin dependent diabetes,21 management of labour,24 discharge planning,29 30 management of intravenous therapy,31 lipid lowering,32 identification of battered women,33 infection control,34 and reduction in rates of falls.32

Five studies assessed the effect of guidelines in terms of the process of care. In four of these, significant changes in at least some of the assessed processes of care were identified.22 23 31 33 For example, the study of Frigoletto et al.31 of the management of labour by nurse midwives found significant differences in practices related to the management of labour between the active management protocol and usual care groups. In the active management group, the frequency of vaginal examinations was significantly higher (mean interval 1.6 hours v 2.5 hours for standard care group); significantly more women received oxytocin to hasten labour (70% v 56%) and significantly fewer women requested and received epidural anaesthesia (54% v 64%). The analysis reported by Larson and Hargiss31 is almost certainly inappropriate, but insufficient data were provided to be certain: there is no explanation as to how the dependent variable was calculated for the analysis of variance (ANOVA), and clustering of the data has probably not been taken into account.

Eight studies measured outcomes of care. In six of these, significant differences in favour of the intervention group were found for at least some outcomes of care.20–22 24 29 30 However, the findings from several of these studies were equivocal, with no significant change being noted in other outcome variables assessed. For some studies, the observed change in what seemed to be the primary outcome measure was non-significant.21 22 30 31

Only one study24 included a formal economic evaluation. This study assessed the effect of medical nutrition therapy for patients with non-insulin dependent diabetes administered according to practice guidelines with medical nutrition therapy administered in line with standard care. Findings were suggestive of a cost efficiency advantage in favour of the control group when one outcome measure (glycated haemoglobin) was considered, but no difference when an alternative outcome measure (fasting plasma glucose concentration) was examined. The studies described by Naylor et al.24 30 evaluating the effects of a protocol for discharge planning compared with standard discharge procedures, and Larson and Hargiss31 evaluating the effectiveness of specially trained nurses who used guidelines in the maintenance of intravenous treatment compared with standard care, suggested that direct healthcare costs are decreased by care driven by a guideline, but these authors did not provide sufficient data to ascertain whether these cost savings were offset by the cost of guideline dissemination and implementation.

SKILL SUBSTITUTION STUDIES
In the remaining six studies (table),18 19 25–28 the performances of nurses operating in accordance with a guideline were compared with standard care, generally provided by a physician. All six studies were randomised controlled trials. The aim was skill substitution, generally in the interests of greater efficiency. Targeted behaviours were: management of dysuria, frequency and vaginal discharge,16 management of low back pain,17 management of headache,27 management of hypertension,18 referral for x ray examination,19 and management of postoperative bleeding after cardiac surgery.20 Study settings were: outpatients,25 26 inpatients,18 the emergency room,28 and primary care.19 28

In these studies, a finding of no significant difference, or of a positive effect in favour of the nurse or other professional allied to medicine, was appropriate. Findings from all six studies which examined the processes and the five which examined outcomes of care generally supported the hypothesis of no difference. However, appropriate sample size calculations were not reported in any study; thus we cannot be sure that there was sufficient power to detect a clinically relevant difference. The finding of no significant difference may therefore have been due to small sample size. Clearly this is
weak evidence, as there was no comparison with patients cared for by nurses not using guidelines.

Discussion

Studies which compared different dissemination and implementation strategies suggest educational interventions are of value in the dissemination of guidelines, and confer a benefit over passive dissemination approaches. This finding is in line with that of Grimshaw and Russell for physicians. However, from the limited evidence presented in the two studies in which there were positive changes in process or outcomes, we cannot tell which forms of education are most effective in bringing about changes in behaviour. Grimshaw and Russell suggested that educational strategies requiring more active participation, such as educational outreach visits and targeted seminars, are more likely to lead to changes in behaviour among physicians; there is an urgent need to investigate whether this assertion holds true for nurses, midwives, and other professions allied to medicine. By contrast, a computerised prompt specific to the patient at the time of consultation did not seem to alter nurses’ behaviour, despite evidence that this is an effective way of changing the behaviour of physicians. However, this finding was based on a small sample of nurses: further research is required before this implementation strategy can be rejected.

Interventions evaluated by Herman et al and by Larson and Hargiss, were complex; complex interventions were also described by French et al. and by Mitchell and Jones. Wensing and Grol have shown that multiple interventions are more likely to be effective in bringing about behavioural change than are single interventions, especially if multiple interventions involve individual instruction, feedback, and reminders. Findings from studies by Herman et al. and Larson and Hargiss provide some evidence of the impact of multiple interventions. However, further research, comparing single and multiple interventions in the same population, is required before we can be sure that the findings of Wensing and Grol are applicable to nursing and professions allied to medicine.

Findings from the 18 studies identified provide some evidence that care driven by guidelines can be effective in changing the process and outcome of care. However, some caution is needed in interpreting the findings from our review, and in generalising these findings to other professions and settings.

Most studies were carried out in a single setting, focused on only one profession, and often involved very few health professionals (in several cases, only one nurse). The possibility that the findings were specific to the setting or to the individual people cannot be ruled out. In some cases, the health professionals who were studied were selected specifically because of their enthusiasm or expertise; the guidelines may have been less effective in the hands of less committed or less highly qualified or experienced staff. As many of the authors themselves recognise, most of the studies were threatened by the Hawthorne effect, as health professionals were aware that their performance was being assessed. The timing of the assessment of process and outcome may also have had an impact: studies we identified did not, in general, provide sufficient information to assess whether any changes in performance found were sustained after the initial impetus of guideline dissemination.

Only three of the 18 studies provided evidence from randomised controlled trials in which the units of randomisation and analysis were the health professional, or from crossover trials where all health professionals experienced each level of intervention. In studies in which the units of randomisation and analysis were the patients, and where the same health professionals provided care for patients in the intervention and control groups, there is the possibility of contamination of the control group by carry over of knowledge of the guideline. Franz et al explicitly recognised that this may have occurred in their study and may be a reason for the lack of significant findings in the primary outcome measures. In controlled before and after studies, bias may be introduced if the baseline characteristics of the sites are different, or if there is some other systematic difference between sites. Shaffer and Wexler recognised that bias may have been introduced by the fact that allocation to the intervention and control groups was not random, but rather was at the discretion of the patients’ regular physicians. In time series designs, there is a risk of contemporaneous changes, a possibility recognised by Tilden and Shepherd.

Many of the studies fall short of the quality criteria for randomised controlled trials, controlled before and after studies, and interrupted time series studies set down by EPOC. Important methodological details were often not reported—for example, few studies specified initially the size of difference in process or outcome that would be considered to be “clinically significant” and reported whether sample sizes were calculated accordingly. Thus we cannot be sure that the studies were adequately powered to detect a clinically important difference. Similarly, in the studies seeking to show no significant difference between care provided by a nurse and driven by a guideline and standard care provided by a physician, appropriate sample size calculations were not reported. In these studies, the finding of no significant difference may have been due to small sample size. Pocock advises that, in general, larger sample sizes are required when the aim is to infer that there is no difference.

Potential sources of bias include a lack of blinding to allocation. If data collectors were aware of which patients were in the intervention and control groups, data collection procedures may have been biased in favour of the intervention group. Few studies reported an intention to treat analysis, although there was often substantial (and, in some cases, differential) attrition from intervention and control samples.
It is mostly impossible to tell whether the guidelines evaluated were based on evidence. Although many were based on a literature review, the extent to which these reviews were systematic was not described, nor were the quality criteria by which any evidence was assessed. The information contained in the papers does not allow us to state whether this lack of an evidence base was due to failure to search for the evidence systematically, or whether there was virtually none. This calls into question the validity of the guidelines.

More research is clearly required in the development, dissemination, implementation, evaluation, and cost effectiveness of clinical guidelines as a strategy for improving professional practice in nursing and professions allied to medicine. Only then will a decision be possible on whether their potential for improving the practice of nurses and professions allied to medicine is as great as it is for doctors.¹

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