Review Article

Reducing Needle Stick Injuries in Healthcare Occupations: An Integrative Review of the Literature

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Needlestick injuries frequently occur among healthcare workers, introducing high risk of bloodborne pathogen infection for surgeons, assistants, and nurses. This systematic review aims to explore the impact of both educational training and safeguard interventions to reduce needlestick injuries. Several databases were searched including MEDLINE, PsycINFO, SCOPUS, CINAHL and Sciencedirect. Studies were selected if the intervention contained a study group and a control group and were published between 2000 and 2010. Of the fourteen studies reviewed, nine evaluated a double-gloving method, one evaluated the effectiveness of blunt needle, and one evaluated a bloodborne pathogen educational training program. Ten studies reported an overall reduction in glove perforations for the intervention group. In conclusion, this review suggests that both safeguard interventions and educational training programs are effective in reducing the risk of having needlestick injuries. However, more studies using a combination of both safeguards and educational interventions in surgical and nonsurgical settings are needed.

1. Introduction

Needle-stick injuries are an important and common occupational injury among healthcare workers. In a UK report, 37% of nurses reported that they have sustained a needle-stick injury at some stage during their career [1]. In Australia, the rate of reported needle-stick injuries is 1 in 5 occupied beds per year which equates to an annual sharps-related injuries incidence of 47,000 [2].

According to the policy of the NHS in the UK, it is compulsory when staff sustain a needle-stick injury to report the incident [3]. However, evidence from the US suggests that more than half of all sharps-related injuries are not reported [1]. Poor reporting of sharps-related injuries reveals a failure to appreciate the potential consequences of such injuries [4]. Rates of detection are also low, for example, only 11% of glove perforations were detected by the physician in a study investigating the use of blunt needles during obstetrical laceration repair surgeries [5].

Needle-stick injuries have been widely recognised as a source of exposure to bloodborne pathogens for workers in healthcare occupations [6]. There are more than 20 bloodborne pathogens that can be transmitted from contaminated needles or sharps, including hepatitis B (HBV), hepatitis C (HCV), and human immunodeficiency virus (HIV; NHS Employees, 2005). The risk of transmission of HIV following a hollow needle injury is approximately 0.3%, compared with 3% for HCV and 30% for HBV [7]. Worldwide, more than 100 healthcare workers have contracted HIV from work-related needle-stick injuries and many thousands have contracted HBV or HCV [2].

Due to the recognised risk of needle-stick injuries, safeguards have been put in place to attempt to lessen the risk of injury. These include the policy of universal precautions and needleless systems to connect with intravenous tubing [5]. Several strategies have been adopted for use in the healthcare setting, including double gloving, having a neutral zone in which to pass sharps, and the use of blunt tip needles [8]. Wearing two pairs of gloves is a practice which protects healthcare workers from patients’ blood and body fluids. A recent study found that in 82% of cases when the outer glove was perforated, the inner gloves had been found to
Despite its benefits, double gloving has not been universally adopted in healthcare settings due to the discomfort and reduced dexterity it may cause. However, studies have shown that double gloving can significantly reduce the risk of needle-stick injuries by preventing puncture wounds and needle perforations. In contrast, single gloving may not provide adequate protection for healthcare workers.

2. Methods

2.1. Design and Search Methods. Using an integrative approach to literature searching, searches were conducted in five different databases, including MEDLINE, PsycINFO, SCOPUS, CINAHL, and Sciencedirect. The terms “glove perforation,” “needle-stick injuries,” “reducing percutaneous injuries,” “glowing tears,” “Reducing sharp injuries,” and “occupational exposure in healthcare” were utilized.

2.2. Search Outcome and Quality Appraisal. Only studies evaluating a needle-stick injury intervention and deliberately designed to contain both a study and a control group within the past 10 years were included. Both studies using new safeguards and providing educational training were reviewed. There were no exclusion criteria regarding the types of participants, duration of intervention, or method of outcome measurement. Studies were rejected if they were not published in a peer-reviewed journal or if the text was not in English. A study was rejected if it did not meet the inclusion criteria determined from the title and abstract during screening.

3. Results

3.1. Study Selection. The search yielded 6942 bibliographic records and fourteen studies met the inclusion criteria [8, 9, 18, 19, 22–24, 26–28] (see Figure 1 for details). No studies were reported in multiple papers.

3.2. Study Characteristics

3.2.1. Study Design. Of the fourteen studies, 10 were randomised control trials of interventions [5, 18–24, 26, 27]. Two studies used a cohort design with no randomisation of participants into conditions [8, 28]. Participants in these cohort studies choose the gloving types at their own discretion. One study was designed in a prospective randomised manner [9]. In this study, the surgeons were randomised to either the study group or the control group by sealed...
envelopes, and, then, the first assistant was automatically allocated into the opposite condition. The remaining study was a quasixperimental design using randomisation procedure for groups rather than for each participant [25]. This study compared outcomes preintervention and postintervention between the intervention and the control group.

3.2.2. Types of Intervention. Of the fourteen studies, ten studies evaluated the use of double gloves in preventing needle-stick injuries [8, 9, 18, 19, 22–24, 26–28]. This included eight studies evaluating the use of double gloves [8, 9, 18, 19, 22, 23, 26, 28], and two studies evaluating variations of double gloving [23, 24, 27]. Three studies evaluated the use of blunt needles in preventing needle-stick injuries [5, 20, 21]. The remaining study compared a bloodborne pathogen educational training with a standard education [25].

Thirteen out of the fourteen studies were conducted in departments of surgery or of emergency medicine and focused on different operations [5, 8, 9, 18–24, 26–28]. Nine of these studies focused on specific surgical procedures, including laparotomy surgery, nonemergent Caesarean delivery, obstetric laceration, elective gastrointestinal procedures, gynaecologic surgery, visceral surgical procedure, arch bar placement for intermaxillary fixation, episiotomy repair after vaginal delivery, orthopaedic and trauma surgery [5, 8, 18–23, 27]. Four studies included general surgeries, which included surgical procedures lasting more than one hour [9, 24, 26, 28]. The remaining study was conducted in a department of nursing [25].

3.3. Study Populations. Thirteen studies recruited participants from hospitals [5, 8, 9, 18–24, 26, 28]. Of these, five recruited surgeons and assistants [5, 9, 21, 24, 28], six only recruited surgeons [8, 19, 20, 22, 23, 27] and two recruited the whole operation team including scrub nurses [18, 26]. The final study recruited students in a department of nursing [25]. The subjects in this study had completed 3 years of academic work and 3 months of clinical practice.

The fourteen studies used different methods to determine sample size. Six studies used individual participants as a unit of measurement, four of these counted “patients treated” as the sample, the sample size varied from 42 to 438 [5, 20, 21, 27]. One study used the number of surgeons who participated in the study as the sample size with 170 surgeons included [19], and the remaining study used the 106 students who participated in the study as the sample size [25]. Four studies used the number of procedures observed as the sample size which varied from 66 to 885 [9, 22–24]. Four studies used the number of pairs of gloves used in the study as the sample size which varied from 300 to 1000 pairs [8, 18, 26, 28]. The follow-up period varied from 2 months to 21 months with five studies not reporting the study duration. Only one study reported the mean age (19 years old) [25]. No study reported the gender proportion of participants.

3.4. Outcome Measures. In terms of outcome measures, the rate of glove perforation was assessed in all studies [5, 8, 9, 18–28]. Other outcome measures included detection rate of glove perforation (5 studies), evaluation of the devices
| Author               | Design                      | Sample criteria                                                                 | Sample size                                                                 | Blinding       |
|---------------------|-----------------------------|---------------------------------------------------------------------------------|------------------------------------------------------------------------------|----------------|
| Punyatanasakchai et al. [18] | Randomized controlled trial | Patients and surgeons from Ramathibodi Hospital, Faculty of Medicine Mahidol University, Bangkok, Thailand | $N = 300 \ (C: n = 150, I: n = 150)$                                        | No blinding   |
| Kovavisarach and Seedadee [19] | Randomized controlled trial | Gynaecologic patients who underwent total abdominal hysterectomy (TAH) with or without bilateral salpingo-oophorectomy (BSO) at Rajavithi Hospital between September 1 1999 to August 31 2000. Primary surgeons and specialist assistants from the same hospital. | $N = 170 \ (C: n = 82, I: n = 88)$                                         | No blinding   |
| Nordkam et al. [20] | Randomized clinical trial   | Patients who underwent laparotomy. Surgeons from the Department of Surgery       | $N = 200 \ (C: n = 100, I: n = 100)$                                        | Not reported   |
| Wilson et al. [5]  | Randomized prospective trial | Patients underwent an obstetric laceration repair in the labor and delivery suite from January 2005 through September 2006. Surgeons from Medical University of South Carolina | $N = 438 \ (C: n = 221, I: n = 217)$                                        | No blinding   |
| Sullivan et al. [21] | Randomized control trial    | All patients requiring nonemergent cesarean deliveries from January to September 2006. Surgeons from Medical University of South Carolina | $N = 194 \ (C: n = 97, I: n = 97)$                                        | No blinding   |
| Caillot et al. [22] | Randomized control trial    | Visceral surgical procedures performed in the surgical emergency department and 5 surgeons in the department. | $N = 100 \ (C: n = 50, I: n = 50)$                                        | Not reported   |
| Gaujac et al. [23] | Comparative randomized trial | Consecutive patients with maxillomandibular fractures. Two emergency room surgeons. | $N = 42 \ (C: n = 47, I: n = 56)$                                        | No blinding   |
| Laine and Aarnio [24] | Randomized prospective trial | All gloves used by the surgeons in 885 surgical operations at Satakunta Central Hospital. | $N = 2462 \ (C: n = 1020, I: n = 1148, combination: n = 294)$ | Not reported   |
| Wang et al. [25]   | Quasiexperimental study     | Students enrolled at Xiang Ya school of Medicine, Department of Nursing, in the 4-year nursing program | $N = 106 \ (C: n = 50, I: n = 56)$                                        | Blinding       |
| Naver and Gottrup [26] | Randomized control study   | Gloves tested on surgeons, assistants and scrub nurses in university hospital, Denmark | $N = 566 \ (C: n = 306, I: n = 260)$                                        | Not reported   |
| Thomas et al. [9]  | Randomized control study    | Surgical procedures lasting more than one hour performed in department of general surgery, lady hardinge medical college and associated srimati sucheta kriplani hospital, New Delhi. | $N = 396 \ (C: n = 198, I: n = 198)$                                        | No blinding   |
| Laine and Aarnio [27] | Randomized control study   | All gloves used in consecutive orthopaedic and trauma operations, conventional and arthroscopic in Satakunta central hospital, Pori, Finland | $N = 972$                                                                  | Not reported   |
| Lancaster and Duff [8] | Cohort study               | Gloves from obstetric and gynaecologic surgical procedures at University of Florida College of Medicine | $N = 100 \ (C: n = 325, I: n = 675)$                                        | No blinding   |
| Na’aya et al. [28] | Cohort study               | Gloves used in general surgical procedures in department of surgery, University of Maiduguri teaching hospital, Nigeria. | $N = 1120 \ (C: n = 240, I: n = 880)$                                        | No blinding   |
| Author                      | Consent rate (CR)/Response rate (RR) | Intervention type | Surgical procedure                  | Randomization procedure and design                                                                 | Study timeline | Outcome measure                                                                 | Results                                                                 |
|-----------------------------|-------------------------------------|-------------------|-------------------------------------|----------------------------------------------------------------------------------------------------|----------------|---------------------------------------------------------------------------------|------------------------------------------------------------------------|
| Punyatañasakchai et al. [18]| Not reported                        | Double gloving   | Episiotomy repair after vaginal delivery | The surgeons were randomly selected one of two envelopes, number 1 representing the single-gloving method and number 2 representing the double-gloving method. | 7 months       | Glove perforation rate, duration of operation, position level of surgeons       | No significant difference in the frequency of perforations between the double-outer gloves (22.6%) and single-gloves (18%). A significant reduction in glove perforations between the double-inner gloves (4.6%) compared with the single-gloves (18%) (P < .05). |
| Kovavisarach and Seedadee [19]| Not reported                        | Double gloving   | Gynaecological surgery             | Primary surgeons were randomly allocated to use either the single-gloving or the double-gloving method. | 12 months      | Glove perforation rate, duration of operation                                   | A significant difference was found in the glove perforation rate between double-inner glove (6.09%) and single gloving group (22.73%). No significant difference between the glove perforation rates in single gloves (22.73%) and in double-inner gloves (19.5%). |
| Nordkam et al. [20]         | Not reported                        | Blunt needle      | Abdominal wall closure             | Surgeons were randomised by envelop to use either blunt needle or sharp needle                      | 6 months       | Glove perforation rate, evaluation of the blunt needle                          | A significantly higher number of surgical procedures with perforations using the sharp needle (P = .003) than with the blunt tapered needle. Detection rate was low (21%). Blunt tapered needles are less convenient |
| Wilson et al. [5]           | Not reported                        | Blunt needle      | Obstetrical laceration repair      | Patients with obstetric lacerations were randomized to repair with either blunt or sharp needles. | 21 months period | Glove perforation rate, evaluation of the blunt needle, and position level of surgeons | No significant difference in the glove perforation rate between blunt and sharp needles. There was poor correlation between reported perforations and those detected by water test. Blunt needles were reported more difficult to use (P = .0001) |
| Sullivan et al. [21]        | Not reported                        | Blunt needle      | Cesarean-delivery closure          | Patients requiring cesarean delivery were assigned randomly to receive closure with either blunt or sharp needles | 21 months      | Glove perforation rate, evaluation of the blunt needle, and duration of operation | A significant reduction in total glove perforation rate for the primary surgeon with blunt needles (7.2%) compared with sharp needles (17.5%) as well as for the assistant surgeons. Poor correlation between reported perforations and those detected by water test. Physicians reported low satisfaction with blunt needles compared with sharp needles (P < .001) |
| Author                          | Consent rate (CR)/Response rate (RR) | Intervention type | Surgical procedure | Randomization procedure and design                                                                 | Study timeline | Outcome measure                                                                 | Results                                                                                   |
|--------------------------------|-------------------------------------|-------------------|-------------------|-----------------------------------------------------------------------------------------------|----------------|---------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------|
| Caillot et al. [22]            | Not reported                        | Double gloving    | Visceral surgical procedures | Visceral surgical procedures performed in the Surgical Emergency Department were randomly assigned to double gloving or single gloving. | 3 months       | Glove perforation rate, detection of the perforation, duration of operation     | Did not adequately compare the rate of glove perforation. Double gloving allowed significantly higher detection rates of glove perforation (P < .001) |
| Gaujac et al. [23]             | Not reported                        | 2 types of double gloving | Arch bar placement | Patients were equally divided into 2 groups. In group 1, 2, sterile surgical gloves were used; in group 2, a nonsterile disposable inner glove was used under a sterile surgical glove. | Not reported  | Glove perforation rate, duration of operation                                   | No significant statistical difference was found between 2 double gloving methods in terms of inner glove perforations |
| Laine and Aarnio [24]          | Not reported                        | Double gloving    | General surgical operations | Patients born in even years were operated on with double gloving and those born in uneven years were operated on with single gloving. | 2 months       | The glove type, the operating time, the type of surgery, the detection rate and location of perforation | A low number of perforations of the inner glove of the double-gloving system were detected. Higher detection of perforation in double-glove system (P < .001). The longer of the operating duration, the higher rate of perforation |
| Wang et al. [25]               | RR: 86%                             | Educational training | NA                | One class was randomly assigned to receive the educational intervention, and the other served as a comparison group receiving standard education. | 4 months       | Changes in knowledge and self-reported universal precautions behaviour, observed adherence to universal precautions, and self-reported needlestick injuries | The group that received the intervention scored significantly higher than the standard education group on both knowledge (P < .001) and behaviour (P = .002), and were less likely to experience needlestick injuries (P = .004) |
| Naver and Gottrup [26]         | Not reported                        | Double gloving    | Various types of gastrointestinal surgery | The surgeons, assistants and scrub nurses were randomized into one of two groups. In group one the operating team was using powder-free single gloves and group two used a powder-free double-gloving system. | Not reported  | Glove perforation rate, detection of the perforation, and the position of the participants | A significant difference between single gloves and inner indicator gloves (P < .005). The surgeon in indicated gloving group obtained high detection rate of glove perforation (P < .0001) |
| Author                | Consent rate (CR)/Response rate (RR) | Intervention type | Surgical procedure       | Randomization procedure and design                                                                 | Study timeline | Outcome measure                                                                 | Results                                                                                                                                                                                                 |
|----------------------|--------------------------------------|-------------------|--------------------------|---------------------------------------------------------------------------------------------------|---------------|---------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Thomas et al. [9]    | Not reported                          | Double gloving    | General surgical operations | The gloving pattern was randomized into two groups of the equal number by sealed envelopes        | Not reported  | Glove perforation rate, detection of the perforation, evaluation of double gloving | In double-gloving pattern, 32 glove perforations were observed, of which 22 were in the outer glove and 10 in the inner glove. Majority of glove perforations (83.3%) went unnoticed. Double gloving was accepted by majority of surgeons. |
| Laine and Aarnio [27]| Not reported                          | Double gloving    | Orthopaedic and trauma surgery | Before the operations, the surgeons were randomised to use either single gloves, double indicator gloves or a combination of two regular surgical gloves on top of each other | 2 months      | Glove perforation rates, detection of perforations, operation types, and duration of operation | Significant difference in perforations of the inner glove in two of indicator gloves and in the regular combination gloves when the outer glove was perforated ($P = .02$) |
| Lancaster and Duff [8]| Not reported                         | Double gloving    | Obstetric and gynecologic surgical procedures | The choice to single versus double glove was left to the discretion of the individual surgeon. | 7 months      | Glove perforation rate, the association between position of the surgeon and perforation rate | 11% of single glove sets contained a perforation whereas only 2% of double glove sets contained a corresponding defect in the inner and outer gloves ($P < .01$) |
| Na’aya et al. [28]   | Not reported                          | Double gloving    | General surgical procedure. | The surgeons wore single or double gloves at their own discretion.                                 | Not reported  | Glove perforation rate, detection rate of the perforation, and duration of operation | A significant greater risk for blood-skin exposure in the single glove sets ($P < .01$) Most perforations were not noticed during the surgery. |
used (4 studies), the relationship between glove perforation and the job level of surgeons (3 studies), the relationship between glove perforation and duration of the operation (8 studies), the relationship between glove perforation and surgical types (3 studies), frequency of glove perforations by position on surgical team (3 studies), changes in knowledge and self-reported universal precautions behaviour, observed adherence to universal precautions, and self-reported needle-stick/sharp injuries (1 study). For the purpose of this review, rate of glove perforation, detection rate of glove perforation, evaluation of devices used, changes in knowledge self-reported universal precautions behaviour, and observed adherence to universal precautions were analysed.

3.5. Study Quality. All fourteen studies contained both an experimental group and a control group. Of the fourteen trials, ten studies reported an adequate randomisation method, all using a cluster randomisation procedure [5, 18–24, 26, 27]. Two cohort studies did not use a randomisation procedure to allocate participants into conditions [8, 28]. Participants in these studies choose the gloving types at their own discretion. In one study, only surgeons were randomised to conditions whereas assistants were automatically allocated into the opposite condition [9]. The remaining study used randomisation procedure for groups rather than for each participant [25].

Outcomes in thirteen of the studies were measured by a water-leak test method, in one study gloves were filled with air then put in water [21]; another used both air and water-leak techniques [5]. Five studies tested unused gloves as controls to test for preexisting minor perforations [5, 9, 18, 19, 21]. One study did not test the glove perforation rate but used questionnaires and direct observation to collect data [25]. Direct observation was used in only three studies excluding the one which did not aim to measure the rate of glove perforation and reported replacement of the perforated gloves during surgery with a similar glove [19, 22, 28]. But only one of the three studies reported that they only used the original gloves as data regardless of any further perforations in the replaced glove [22]. Only one study reported that the individual testing of the gloves was blinded to the allocation of the glove wearer [25].

One study included blinding of participants to condition [25], eight studies did not include blinding participants [5, 8, 9, 18, 19, 21, 23, 28], and five studies did not report blinding of participants to condition [20, 22, 24, 26, 27]. None of the studies reported participant refusal rates and withdrawal rates.

Selection issues were a potential source of bias for a number of studies. Thirteen of the fourteen studies were conducted in departments of surgery or emergency medicine [5, 8, 9, 18–24, 26–28], and nine studies focused on different surgical procedures [5, 8, 18–23, 27]. Only four studies evaluated interventions being used in different hospital units across different types of operations [9, 24, 26, 28], and only one study was conducted in a department of nursing [25]. The use of only surgical procedures included in these studies, and the homogeneity of individual participants may be a potential bias. Only two studies recruited the whole operation team including the principal surgeon, the surgical assistants, and the scrub nurses [18, 26]; eleven studies included either surgeons alone or both surgeon and first assistants [5, 8, 9, 18–24, 26, 28]. Because most of the studies tested the intervention of double gloving and blunt needles, gloves were collected immediately after each surgery. Only one study that was conducted in a department of nursing reported the consent rate and the dropout rate (86%) [25].

The studies differed in the method of comparing intervention and control groups which may bias interpretation of the results. Of the thirteen studies which aimed to compare the perforation rates [5, 8, 9, 18–24, 26–28], seven compared the gloves in the single-gloving condition separately with both the outer and inner gloves used in double-gloving conditions [5, 8, 18, 19, 21, 26, 28], two compared the inner gloves in both conditions [23, 27], two compared the total perforation numbers in both conditions [9, 24], one compared the number of surgical procedures with perforations [20], and one study did not report the method used [22]. Also the studies differed in a variety of ways, including the location of the intervention (country, hospital units), the time period of the study, and the number of participants. Four studies did not report the number of surgeons in the study instead reported the number of patients [5, 19–21], so it is possible that only a few surgeons generated most of the data which could limit the generalisability of the results.

3.6. Outcomes. The effectiveness of interventions using new devices to reduce needle-stick injuries in healthcare occupations was mainly defined by comparing the numbers of glove perforations or numbers of needle-stick injuries from the study populations with those of control populations. The effectiveness of educational training to reduce needle-stick injuries in healthcare occupations focused on comparing the changes in needle-stick injuries knowledge between study and control populations.

3.7. Double Gloving. Within the ten studies which evaluated double gloving or combinations of gloving [8, 9, 18, 19, 22–24, 26–28], seven compared single and double gloving [8, 9, 18, 19, 22–24, 26–28], two compared single, double, and combination gloving [23, 24, 27], and one compared one double-gloving method [23]. Eight studies reported an overall reduction by 9 to 15% in glove perforations in inner gloves under double-gloving conditions compared to those under single gloving conditions [8, 9, 18, 19, 22–24, 26–28]. One study only used descriptive statistics rather than inferential analysis to measure the differences of glove perforation rate [22]. One study, comparing two double-gloving methods with one, used two sterile surgical gloves and one used a nonsterile disposable inner glove under a sterile surgical glove [23], but did not find any significant statistical differences between groups.

Of the four studies who reported the detection rate of glove perforation [9, 22, 24, 28], two reported a low detection rate in the double-gloving method [9, 20] and two reported a high detection rate when double gloving was used [24, 26]. The two studies which demonstrated a high
detection rate differed from the two studies that found a low detection rate, in the brand of gloves used, which may be a factor in interpreting the results [9, 22, 24]. In these gloves, glove perforation during surgery results in an inflow of fluid between the two pairs of gloves. The wet area of the inner glove then appears as a bright green spot under the perforation area of the outer glove, which can be easily noticed by the wearer [24].

3.8. Blunt Needles. Three of the fourteen studies reviewed in this paper evaluated the use of blunt needles in reducing needle-stick injuries [5, 20, 21]. All of these studies were randomised control trials. Two studies reported a significant reduction by 9% to 16% in glove perforations for outer gloves of double gloving [20, 21], whereas one study did not find a significant difference in glove perforation between using blunt and sharp needles [5]. That study compared the rate of glove perforation for blunt and sharp needles used during obstetrical laceration repair which are said to require less time to complete [5], which may explain the apparent anomaly. The remaining two studies found a significant reduction in glove perforation when blunt needles were used during laparotomy [20] and Caesarean delivery [21]. The three studies reported the detection rate of glove perforation [5, 20, 21]. All of the three studies reported by surgeons that blunt needles were less convenient to use and associated with less satisfaction.

3.9. Educational Training. This quasiexperimental study examined the impact of structured training on prevention of occupational exposure to bloodborne pathogens on knowledge, behaviour, and incidence of needle-stick injuries among student nurses [25]. It reported a significantly higher score on both knowledge ($P < .001$) and behaviour ($P = .002$) in the group who received the bloodborne pathogens training. The self-reported needle-stick injuries were significantly lower for the bloodborne pathogens training group though they were not observed to practice universal precautions significantly more frequently than those in the control group [25].

4. Discussion

4.1. Main Findings. The reviewed interventions on needle-stick injuries includes components of double gloving, using blunt needles and educational training, and demonstrates there are significant reductions in needle-stick injuries following interventions as measured by glove perforations and changes in bloodborne pathogen knowledge. Results showed that interventions that use safeguard devices (double gloving and blunt needles) lead to a reduction in needle-stick injuries among healthcare workers. Knowledge regarding bloodborne pathogens was a major outcome measure in just one study [25] and results suggest that a structured bloodborne pathogen educational training program can lead to improvements in knowledge and a reduction in self-reported needle-stick injuries [25]. As such, the limited evidence regarding the effectiveness of educational training on bloodborne pathogen knowledge and behaviour was inconclusive.

The results of this review are consistent with a much earlier review of interventions to prevent needle-stick injuries [6]. In this study, a reduction in the number of glove perforations was found in eight out of eleven studies. The current review included more recent studies in this field with more flexible study design (any case control studies) and types of interventions (double gloving, blunt needles and educational training). The current research findings are useful not only for encouraging healthcare workers to use double gloves and blunt needles during operations, but also for attracting policy makers to promote the universality of safeguards.

4.2. Limitation of the Reviewed Studies. The fourteen studies differed in a variety of ways, including the surgical procedures, the type of intervention, the location of the intervention (country, hospital units), the time period of the study, and the number of participants. Most studies reported relatively serious methodological flaws such as study procedure (13 studies), randomisation methods (5 studies), and statistical tests (10 studies). In addition, thirteen studies failed to report drop-out rate, blinding procedures, consent rate, and exact number of participants which may bias results [8, 9, 18, 19, 22–24, 26–28]. In the study conducted in a department of nursing, the possibility of communication between the study group and the control group may have influenced the effect of the educational training [25]. The lack of blinding may have introduced bias into the study as the participants in both groups knew the aim of the study and as a result may have paid more attention or been more careful during operations, which may decrease the function of the control group. Thus, caution about the differences within the fourteen studies is needed when drawing robust conclusions based on the results.

Of the fourteen studies, only one examined the impact of a structured educational training program on prevention of occupational exposure to bloodborne pathogens on knowledge, and behaviour among student nurses [25]. Although the other studies which investigated the effectiveness of safeguard devices have obtained significant results, a number of studies on the prediction of behaviours in health fields using social cognitive models suggest that intentions are the immediate antecedent to performing a specific behaviour [29]. In general, the stronger the intention to engage in behaviour is, the more likely it will be performed [29]. It is thought that if certain behaviours are planned to be performed in specified conditions (e.g., “I will wear double gloves and perform very carefully during the surgical procedure”) and are consciously prepared (e.g., “set alarm”), when conditions are encountered the cues stimulate automatic activation of the behaviour [30]. In line with this evidence, it may be that a more effective intervention would contain structured educational training aimed at changing healthcare workers’ attitudes and intentions to prevent needle-stick injuries, or combine educational training with a safeguards intervention, using implementation intentions.

All the reviewed studies in this paper contained an experimental group and a control group, and most had
proper randomised controlled procedures. However, it may raise ethical issues if interventions are designed in a randomised controlled manner which may put the healthcare workers at a high risk of incurring needle-stick injuries in the control group. The healthcare workers in the control group may have increased risk of experiencing needle-stick injuries compared to the healthcare workers randomised to the experimental group. Thus, such possible ethical issues need to be considered in order to minimise the risk for healthcare workers.

5. Conclusions

More studies are needed to evaluate interventions in nonsurgical settings, such as departments of nursing and other hospital units and among other healthcare personnel such as nurses. The current evidence suggests that both safeguard interventions and educational training programs are effective in reducing the risk of having needle-stick injuries. However, there are insufficient studies using a combination of both safeguards and educational interventions in surgical and nonsurgical settings. In future research, evaluations of these two types of interventions in both randomised controlled trials and in studies utilising other designs are needed.

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