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Early postnatal discharge from hospital for healthy mothers and term infants

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[Intervention Review]

Early postnatal discharge from hospital for healthy mothers and term infants

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ABSTRACT

Background

Length of postnatal hospital stay has declined dramatically in the past 50 years. There is ongoing controversy about whether staying less time in hospital is harmful or beneficial. This is an update of a Cochrane Review first published in 2002, and previously updated in 2009.

Objectives

To assess the effects of a policy of early postnatal discharge from hospital for healthy mothers and term infants in terms of important maternal, infant and paternal health and related outcomes.

Search methods

We searched the Cochrane Pregnancy and Childbirth Group's Trials Register, ClinicalTrials.gov, the WHO International Clinical Trials Registry Platform (ICTRP) (21 May 2021) and the reference lists of retrieved articles.

Selection criteria

Randomised controlled trials comparing early discharge from hospital of healthy mothers and term infants (at least 37 weeks' gestation and greater than or equal to 2500 g), with the standard care in the respective settings in which trials were conducted. Trials using allocation methods that were not truly random (e.g. based on patient number or day of the week), trials with a cluster-randomisation design and trials published only in abstract form were also eligible for inclusion.

Data collection and analysis

Two review authors independently assessed trials for inclusion and risk of bias, extracted and checked data for accuracy, and assessed the certainty of evidence using the GRADE approach. We contacted authors of ongoing trials for additional information.

Main results

We identified 17 trials (involving 9409 women) that met our inclusion criteria. We did not identify any trials from low-income countries. There was substantial variation in the definition of 'early discharge', ranging from six hours to four to five days. The extent of antenatal preparation and midwifery home care offered to women following discharge in intervention and control groups also varied considerably among trials. Nine trials recruited and randomised women in pregnancy, seven trials randomised women following childbirth and one did not report whether randomisation took place before or after childbirth.



Risk of bias was generally unclear in most domains due to insufficient reporting of trial methods. The certainty of evidence is moderate to low and the reasons for downgrading were high or unclear risk of bias, imprecision (low numbers of events or wide 95% confidence intervals (CI)), and inconsistency (heterogeneity in direction and size of effect).

Infant outcomes

Early discharge probably slightly increases the number of infants readmitted within 28 days for neonatal morbidity (including jaundice, dehydration, infections) (risk ratio (RR) 1.59, 95% CI 1.27 to 1.98; 6918 infants; 10 studies; moderate-certainty evidence). In the early discharge group, the risk of infant readmission was 69 per 1000 infants compared to 43 per 1000 infants in the standard care group. It is uncertain whether early discharge has any effect on the risk of infant mortality within 28 days (RR 0.39, 95% CI 0.04 to 3.74; 4882 infants; two studies; low-certainty evidence). Early postnatal discharge probably makes little to no difference in the number of infants having at least one unscheduled medical consultation or contact with health professionals within the first four weeks after birth (RR 0.88, 95% CI 0.67 to 1.16; 639 infants; four studies; moderate-certainty evidence).

Maternal outcomes

Early discharge probably results in little to no difference in women readmitted within six weeks postpartum for complications related to childbirth (RR 1.12, 95% CI 0.82 to 1.54; 6992 women; 11 studies; moderate-certainty evidence) but the wide 95% CI indicates the possibility that the true effect is either an increase or a reduction in risk. Similarly, early discharge may result in little to no difference in the risk of depression within six months postpartum (RR 0.80, 95% CI 0.46 to 1.42; 4333 women; five studies; low-certainty evidence) but the wide 95% CI suggests the possibility that the true effect is either an increase or a reduction in risk.

Early discharge probably results in little to no difference in women breastfeeding at six weeks postpartum (RR 1.04, 95% CI 0.96 to 1.13; 7156 women; 10 studies; moderate-certainty evidence) or in the number of women having at least one unscheduled medical consultation or contact with health professionals (RR 0.72, 95% CI 0.43 to 1.20; 464 women; two studies; moderate-certainty evidence).

Maternal mortality within six weeks postpartum was not reported in any of the studies.

Costs

Early discharge may slightly reduce the costs of hospital care in the period immediately following the birth up to the time of discharge (lowcertainty evidence; data not pooled) but it may result in little to no difference in costs of postnatal care following discharge from hospital, in the period up to six weeks after the birth (low-certainty evidence; data not pooled).

Authors' conclusions

The definition of 'early discharge' varied considerably among trials, which made interpretation of results challenging. Early discharge probably leads to a higher risk of infant readmission within 28 days of birth, but probably makes little to no difference to the risk of maternal readmission within six weeks postpartum. We are uncertain if early discharge has any effect on the risk of infant or maternal mortality. With regard to maternal depression, breastfeeding, the number of contacts with health professionals, and costs of care, there may be little to no difference between early discharge and standard discharge but further trials measuring these outcomes are needed in order to enhance the level of certainty of the evidence. Large well-designed trials of early discharge policies, incorporating process evaluation and using standardized approaches to outcome assessment, are needed to assess the uptake of co-interventions. Since none of the evidence presented here comes from low-income countries, where infant and maternal mortality may be higher, it is important to conduct future trials in low-income settings.

PLAIN LANGUAGE SUMMARY

Early postnatal discharge from hospital for healthy mothers and term infants

We set out to determine from randomised controlled trials the effects of a policy of early postnatal discharge from hospital for healthy mothers and term infants (born at 37 weeks of pregnancy or later) on maternal, infant and paternal health and related outcomes.

What is the issue?

Problems can develop or become clear after the birth of a baby. For example, women can experience excessive bleeding and infections, have problems with initiating breast feeding, and lack confidence in the care of their infants, and the baby may not thrive. In years gone by, women were kept in hospital to prevent or deal with these issues. The length of time women spend in hospital after childbirth has fallen dramatically in many countries over the past 50 years.

Why is this important?

It is not known whether having a shorter stay in hospital after birth is beneficial or harmful to women and their newborn infants. Earlier discharge of mothers and their babies has potential advantages, including a familiar environment and better sleep, less exposure to artificial schedules imposed in the hospital environment and decreased exposure to infection risks. However, leaving hospital earlier



may result in missed opportunities for breastfeeding and infant care support and identification of infant and maternal health problems following birth. This review of trials compared the policy of early discharge after childbirth with standard length of stay and care at the time of the study.

What evidence did we find?

We searched for evidence in May 2021 and identified 17 trials involving 9409 women. The evidence is of low to moderate certainty because of limitations in the ways the studies were conducted. There was considerable variation in how early discharge was defined, ranging from six hours to four to five days. In most of the trials included in this review, early discharge was accompanied by some level of nursing or midwifery support. None of the trials took place in low-income countries.

Early discharge probably slightly increases the number of babies readmitted to hospital within 28 days of being born (10 studies, 6918 babies, moderate-certainty evidence). It is uncertain whether early discharge has any effect on the risk of babies dying within 28 days (two studies, 4882 babies). Early postnatal discharge probably makes little to no difference to the number of babies having at least one unscheduled medical consultation or contact with health professionals within the first four weeks after birth (four studies, 639 babies, moderate-certainty evidence).

Early discharge probably results in little to no difference in the number of women readmitted to hospital within six weeks of giving birth for complications related to childbirth (11 studies, 6992 women, moderate-certainty evidence). No deaths were reported. The number of women having at least one unscheduled medical consultation or contact with health professionals was not clearly different (two studies, 464 women, moderate-certainty evidence). Similarly, early discharge may result in little to no difference in the risk of depression within six months after giving birth (five studies, 4333 women, low-certainty evidence).

Early discharge probably results in little to no difference in the number of women breastfeeding at six weeks after giving birth or in the number of women having at least one unscheduled medical consultation or contact with health professionals.

Early discharge may slightly reduce the costs of hospital care with little to no difference in the cost of care from discharge to six weeks after the birth.

What does this mean?

The risk of babies being readmitted to hospital is probably higher following early discharge of mothers and their babies from hospital after the birth, but is probably not higher for women being readmitted to hospital after early discharge. We are uncertain about the risk of death for babies and mothers following early discharge, because these are uncommon events. Differences between early discharge and standard discharge in terms of maternal depression, breastfeeding, the number of contacts with health professionals and costs of care are not clearly different, and until further studies are done to investigate these factors, the evidence remains of low certainty.

SUMMARY OF FINDINGS

Summary of findings 1. Early discharge compared to standard discharge for healthy mothers and term infants

Early discharge compared to standard discharge for healthy mothers and term infants

Patient or population: healthy mothers and term infants Setting: maternity units

Intervention: early discharge

Comparison: standard discharge

Outcomes	Anticipated absolute effects [*] (95% CI)		Relative effect (95% CI)	№ of partici- pants	Certainty of the evidence	Comments
	Risk with standard discharge	Risk with early dis- charge	_ (5570 CI)	(studies)	(GRADE)	
Infants readmitted for neonatal morbid- ity within 28 days	Study population		RR 1.59 (1.27 to 1.98)	6918 (10 RCTs)	⊕⊕⊕⊝ MODERATE ¹	Removing four trials with a high
ity within 28 days	43 per 1000	68 per 1000 (54 to 85)		(10 KC1S)	MODERATE -	level of missing data in sensitiv- ity analysis sug- gests that it is un- certain whether early discharge, compared with standard dis- charge, has any effect on the risk of infant read- mission with- in 28 days (RR 1.10, 95% CI 0.67 to 1.81; 3647 in- fants).
Infant mortality within 1 year	Study population		RR 0.45 (0.07 to 2.77)	4984 (3 RCTs)	⊕⊕⊝⊝ LOW 2 3	
	1 per 1000	1 per 1000 (0 to 3)	()	()	2010	
Number of infants having contact with healthcare professionals regarding in-	Study population		RR 0.88 (0.67 to 1.16)	639 (3 RCTs)	⊕⊕⊕⊙ MODERATE ⁴	
fant health issues within 4 weeks of birth	242 per 1000	213 per 1000 (162 to 281)	(0.01 (0 1.10)			



4

Farlyr	Women readmitted within 6 weeks	Study population	RR 1.12 (0.82 to 1.54)	6992 (11 RCTs)	⊕⊕⊕⊝ MODERATE ⁴
octnatal		25 per 1000 28 per 1000 (21 to 39)	(0.02 to 1.0 t)	(III Kers)	
Early postnatal discharge from hospital for healthy mothers and term infants (Review)	Maternal mortality within 6 weeks - not reported		-	-	-
from h	Women probably depressed within 6 months	Study population	RR 0.80 (0.46 to 1.42)	4333 (5 RCTs)	⊕⊕⊝⊝ LOW 4 5
osnital fo	lionais	437 per 1000 349 per 1000 (201 to 620)	(0.10 (0 1.12)	(5 (6 (5)	
	Women breastfeeding (exclusively or partially) at 6 weeks postpartum	Study population	RR 1.04 (0.96 to 1.13)	7156 (10 RCTs)	⊕⊕⊕⊝ MODERATE ⁶
hv moth	partially at o weeks postpartan	644 per 1000 670 per 1000 (619 to 728)	(0.30 to 1.13)	(10 (10))	
N	Number of women having contact with healthcare professionals regarding ma- ternal health issues within 6 weeks of giving birth	Study population	RR 0.72 - (0.43 to 1.20)	464 (2 RCTs)	⊕⊕⊕⊝ MODERATE ⁴
torm info		133 per 1000 96 per 1000 (57 to 160)	(0.45 (0 1.20)	(21(013)	MODERATE '
sto (Doutions)	Costs of hospital care in the period im- mediately following the birth up to the time of discharge	3 studies reported lower hospital care costs for women in the early discharge groups compared to standard care group but the data were not suitable for analysis.	-	1011 (3 RCTs)	⊕⊕⊙© LOW 7
	Costs of postnatal care following dis- charge from hospital in the period up to 6 weeks after the birth	One study (459 women) reported higher postna- tal care costs in the early discharge group. Anoth- er 2 studies (552 women) reported slightly lower postnatal care costs in the early discharge group.	-	1011 (3 RCTs)	⊕⊕⊙© LOW 7

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*The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval; RCT: Randomised controlled trial; RR: Risk ratio

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

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¹Downgraded one level for risk of bias: some domains at high risk due to inadequate random sequence generation, lack of blinding of outcome assessors and differential attrition

²Downgraded one level for risk of bias: high or unclear risk of bias in random sequence generation, high risk of attrition bias

³Downgraded one level due to imprecision: very few events

⁴Downgraded one level due to imprecision: 95% confidence interval is consistent with possible benefit and possible harm

⁵Downgraded one level due to inconsistency: heterogeneity in direction and size of effect

⁶Downgraded half a level for risk of bias (unclear risk for random sequence generation, allocation concealment and selective reporting) and half a level for inconsistency (unexplained statistical heterogeneity)

⁷Downgraded two levels for imprecision: no overall effect estimate and no confidence intervals



BACKGROUND

Since the 1970s, and earlier in some Western countries, there has been a steady decline in the length of time mothers spend in hospital after giving birth. From a standard hospital lying-in period of between eight to 14 days in the 1950s (Rush 1989), length of postnatal hospital stay for an uncomplicated vaginal birth in Australia, Canada, Sweden and the USA is now around two to three days or less (Campbell 2016). Within the UK context, postnatal hospital stay has been decreasing since the 1970s (NHS Digital 2019). In 1975, 42% of postnatal women in England remained in hospital for seven days or more after birth, and in 2018-19 this proportion had declined to 2%. Currently, 20% of women go home the same day of birth, with a further 38% on day one and 23% on day two, making a total of 82% being discharged within three days of birth. In parts of the USA, hospital stays of 12 to 24 hours for uncomplicated vaginal births, and 48 to 72 hours for uncomplicated caesarean births, had become standard by the mid-1990s (Braveman 1995; Declerq 1997). Concern about possible adverse outcomes of early discharge led the USA Congress to pass legislation in 1996 mandating that private insurers cover postnatal stays of at least 48 hours after a vaginal birth, and 96 hours after a caesarean section. However, four years later, the majority of newborn term (at least 37 weeks' gestation) infants were still being discharged 'early' (Lansky 2006).

Description of the condition

The standard length of time a woman stays in hospital after giving birth varies between countries. In Australia, a five to seven day stay was the norm until the mid-1990s (Day 1997), but in most other Western countries a shorter length of postnatal hospital stay had become standard well before this time (Braveman 1995; Rush 1989). As a result of these variations in practice, a length of stay of less than three days postpartum, which would have been considered standard in the late 1980s in countries such as the UK and the USA, would have been considered 'early discharge' in a country like Australia, at the same time point in history. This variation is reflected in the published literature on early discharge, with study participants discharged as early as six to 12 hours postpartum and as late as three to four days after the birth being considered in the early discharge category (Boulvain 2004; Yanover 1976).

Traditionally, women have been kept in hospital to prevent a number of possible adverse outcomes, which include delays in detecting and treating infant and maternal morbidity, greater occurrence of breastfeeding problems leading to earlier weaning, decreased maternal confidence due to lack of professional support, less maternal satisfaction with postnatal hospital care, higher prevalence of maternal depression, and increased infant and maternal readmissions (Braveman 1995; Britton 1994).

A series of population-based cohort studies and case-control studies have been conducted to assess the relationship between length of postnatal hospital stay and maternal and infant health-related outcomes. These studies have produced conflicting results. Two studies exploring breastfeeding initiation and duration in Australia and Sweden found no association (Brown 2004; Waldenström 2004), whereas a study in California, USA found that women who left hospital earlier than the standard length of stay of two nights for a vaginal birth, and four nights for a caesarean

section, were at a slightly increased risk of ceasing breastfeeding earlier (Heck 2003).

Studies examining the association between length of stay and hospital infant readmissions also report conflicting results. Liu 1997, Liu 2000 and Danielson 2000 found an association between shorter length of postnatal stay and increased rates of hospital readmission. In contrast, Edmonson 1997, Kotagol 1999 and Mandl 1997 reported no evidence of an association between shorter hospital stay and readmissions. The most recent study by Harron 2017 and colleagues found no association between 34 and 36 weeks gestation, each additional day of postnatal stay in hospital was associated with a 8.6% decreased risk of readmission (95% confidence interval (CI) 6.1% to 10.5%). Another study conducted by Lain and colleagues found an association between longer length of birth hospitalisation and risk of readmission (Lain 2014).

Several studies have also examined the relationship between postnatal length of stay and jaundice-related readmissions. The most recent population-based study found that infants discharged from hospital in the first two days after birth were more likely to be readmitted for jaundice than infants who had a postnatal length of stay of three days or more (Lain 2015). Older studies raised concerns about early discharge leading to an increase in the number of infants developing severe hyperbilirubinaemia (Catz 1995; Maisels 1995). However, each of these reports was based on a very small number of cases, with no adequate comparison group for assessing the contribution early discharge may have made. Another large retrospective cohort study in the state of Washington, USA using routinely collected data for births between 1989 and 1990 found that newborns discharged before 30 hours of age had a significantly higher rate of mortality in the first month of life, and in the first year of life, than those newborns who stayed in hospital longer (Malkin 2000).

One important factor, which makes comparisons between studies conducted in different countries problematic, is the extent to which earlier discharge is accompanied by co-interventions (for example, varying levels of antenatal preparation and postdischarge support). An early discharge programme that includes frequent home visits from a health professional is likely to produce different results to an early discharge policy that does not include any after care in the community. The level of primary care support available to postnatal women once they leave hospital irrespective of the timing of discharge varies considerably between countries. In the UK, access to medical care from general practitioners and community midwives is universal and free at the point of service, and whilst the number of postnatal contacts in the community has decreased from seven to four over the last 30 years (Care Quality Commisson 2017; MacArthur 2002), community postnatal care provision is still deemed a critical aspect of postnatal care. This high level of service provision is possibly the reason there has been much less concern about the impact of earlier discharge of mothers and babies in the UK than in countries where access to primary care after discharge from hospital is limited (World Health Organization 2013). A population-based study of postnatal care provided to healthy newborns conducted in 19 US states found that 11% to 49% of newborn infants discharged 'early' did not receive a follow-up home visit within one week postpartum (Lansky 2006).

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Description of the intervention

Shorter length of stay has been promoted in midwifery-led care settings (for women and infants at low risk of complications) such as birth centres and is consistent with a move away from an illness orientation in maternity care towards a more familycentred approach (Rush 1989; Waldenström 1987). From this perspective, earlier discharge of mothers and babies affords many potential advantages. These include the opportunity for all family members to be together as they get to know the baby, which would contribute to improved bonding, greater involvement of the father and less sibling rivalry (Britton 1994); the possibility that mothers may obtain more rest and sleep in their own home environments, where they are not exposed to constant interruptions and noise associated with hospital routines (Rush 1989); decreased exposure of the mother and the infant to nosocomial infections (Hellman 1962); enhanced maternal confidence in caring for the baby in the home environment (Rush 1989); and potentially fewer breastfeeding problems due to less conflicting advice and less exposure of the infant to the artificial schedules imposed in the hospital environment (Hellman 1962). The potential advantages of shorter postnatal hospital stays are in many respects the mirror image of the adverse outcomes identified by those with concerns for the safety of mothers and babies. There have been concerns that early discharge from hospital leaves insufficient time to establishing breastfeeding with direct support from health care professionals, resulting in an increase in feeding-related problems such as weight loss, dehydration and jaundice (Gupta 2006). Concerns have also been raised that in the absence of further community support, early discharge may increase the delay in identification and treatment of maternal and infant morbidity including cardiac defects, as well as vitamin K administration and provision of contraceptive advice.

How the intervention might work

This review will look at the evidence comparing a policy of early discharge following birth with a standard length of stay in hospital. There is uncertainty whether the length of stay may be of benefit or otherwise to women and their families. While staying in hospital under the direct supervision of healthcare professionals may prevent potential morbidities and reduce re-admission rates, earlier discharge home may mean more positive experiences of the first few days postpartum, improved bonding and higher breastfeeding rates.

Why it is important to do this review

While there are several other published reviews on this topic, the continuing reduction of length of postnatal stay in a number of countries and absence of clear evidence regarding safety, potential benefits for families, and costs associated with earlier postnatal discharge warrant the publication of a systematic review assessing current evidence from randomised controlled trials (RCTs).

Several systematic reviews of early postnatal discharge have been published (Beck 1991; Benahmed 2017; Braveman 1995; CETS 1997; Grullon 1997; Jones 2020; Margolis 1995; Norr 1987). Due to the lack of trial evidence in this area of study, the most recent of these have reviewed data from randomised controlled trials and non-randomised studies to consider the effects on neonatal and maternal morbidity, maternal and infant readmissions and women's views of care (Benahmed 2017; Jones 2020). Whilst Benahmed and colleagues concluded that a link between early discharge and neonatal and maternal morbidity could neither be confirmed nor ruled out (Benahmed 2017), Jones and colleagues found that infants discharged early after birth were more likely to be admitted within 28 days. Furthermore, evidence from meta-analysed interrupted times series data showed that the introduction of postnatal minimum length of stay policies in the United States between 1990-2000 was associated with a long-term reduction in neonatal hospital readmission rates (Jones 2020).

Due to the paucity of published trials in the review's earlier version, there was inadequate statistical power to detect increases in maternal or infant readmissions or rarer mortality outcomes (Brown 2002). Since the original review, new trials have been conducted which will contribute to the evidence base.

OBJECTIVES

To assess the effects of a policy of early postnatal discharge from hospital for healthy mothers and term infants in terms of important maternal, infant and paternal health, and related outcomes.

METHODS

Criteria for considering studies for this review

Types of studies

All trials in which women or caregivers or institutions are allocated randomly or quasi-randomly (e.g. alternate allocation) to different policies in relation to the timing of postnatal discharge of healthy mothers and term infants. Trials with a cluster-randomisation design were eligible for inclusion. Studies with a cross-over design would not be appropriate for this clinical question, so we did not include them. We included all RCTs regardless of publication status.

Types of participants

Women who give birth in hospital to a healthy infant of at least 2500 g at term (37 to 42 weeks) who are deemed eligible for 'early discharge'.

Types of interventions

A policy of early postnatal discharge from hospital for healthy mothers and infants born at term where 'early discharge' refers to discharge that is earlier than standard care in the setting in which the intervention is implemented. Standard care refers to the normal postnatal discharge policy in the setting in which the trial is implemented.

Types of outcome measures

Primary outcomes

Infant outcomes

 Infants readmitted for neonatal morbidity (including jaundice, dehydration, infections) within seven days, and within 28 days after birth.

Maternal outcomes

 Women readmitted for complications related to childbirth (including postpartum haemorrhage, retained products of conception, infection, postpartum psychosis) in the first six weeks after giving birth.



- Women scoring above the cut-off score indicating probable depression on a well-validated standardized instrument for measuring depression within six months after giving birth.
- Women breastfeeding (exclusively or partially) at six weeks, 12 weeks and six months after giving birth.

Secondary outcomes

Infant outcomes

- Infant mortality within 28 days and within one year after birth.
- Duration of infant readmissions for infants readmitted within seven days, and within the first 28 days after the birth.
- Total duration of infant hospitalisation over the first 28 days.
- Infants attending hospital casualty or emergency department within seven days, and the first six weeks after birth.
- Number of contacts with health professionals regarding infant health issues within seven days, and the first six weeks after birth.

Maternal outcomes

- Maternal mortality within six weeks and within one year after giving birth.
- Duration of readmissions for women readmitted within first six weeks after giving birth.
- Total duration of maternal hospitalisation over the first six weeks after giving birth (including period immediately after giving birth in addition to any readmissions).
- Women attending hospital casualty or emergency department within first six weeks after giving birth.
- Number of contacts with health professionals regarding maternal health issues within the first six weeks after giving birth.
- Women reporting tiredness or exhaustion in the first six weeks after giving birth.
- Women reporting physical health problems (including perineal pain, perineal infection, breast soreness, breast infection, caesarean wound pain, caesarean wound infection) in the first six weeks after giving birth.
- Women reporting that they lacked confidence about caring for their baby in the first month and the first six months after being discharged from hospital.
- Women reporting infant feeding problems in the first six weeks after giving birth.
- Women reporting they received conflicting advice regarding breastfeeding in the first six weeks giving birth.
- Women satisfied with overall postnatal care (including in hospital and/or post-discharge care).
- Women who perceive their length of hospital stay as too short.
- Women who perceive their length of hospital stay as too long.

Partner outcomes

- Partners reporting that they lacked confidence about caring for their baby in the first month and the first six months after the baby came home from hospital.
- Partners reporting a high level of involvement with their baby in the first month and in the first six months after the birth.

Economic outcomes

- Costs of hospital care in the period immediately following the birth up to the time of discharge.
- Costs of postnatal care following discharge from hospital in the period up to six weeks after the birth, including community midwife, lactation consultant, general practice, specialist and outpatient visits; readmissions to hospital; attendances at day-stay programs; in-patient stays in mother and baby units.
- Costs of practical support following discharge from hospital in the period up to six weeks after the birth, including paid and unpaid home help, care of the baby and of siblings.

Search methods for identification of studies

The following methods section of this review is based on a standard template used by Cochrane Pregnancy and Childbirth.

Electronic searches

For this update, we searched Cochrane Pregnancy and Childbirth's Trials Register by contacting their Information Specialist (21 May 2021).

The Register is a database containing over 25,000 reports of controlled trials in the field of pregnancy and childbirth. It represents over 30 years of searching. For full current search methods used to populate Pregnancy and Childbirth's Trials Register including the detailed search strategies for CENTRAL, MEDLINE, Embase and CINAHL; the list of handsearched journals and conference proceedings, and the list of journals reviewed via the current awareness service, please follow this link.

Briefly, Cochrane Pregnancy and Childbirth's Trials Register is maintained by their Information Specialist and contains trials identified from:

- 1. monthly searches of the Cochrane Central Register of Controlled Trials (CENTRAL);
- 2. weekly searches of MEDLINE (Ovid);
- 3. weekly searches of Embase (Ovid);
- 4. monthly searches of CINAHL (EBSCO);
- 5. handsearches of 30 journals and the proceedings of major conferences;
- 6. weekly current awareness alerts for a further 44 journals plus monthly BioMed Central email alerts.

Search results were screened by two people and the full text of all relevant trial reports identified through the searching activities described above were reviewed. Based on the intervention described, each trial report was assigned a number that corresponded to a specific Pregnancy and Childbirth review topic (or topics), and was then added to the Register. The Information Specialist searched the Register for each review using this topic number rather than keywords. This resulted in a more specific search set that has been fully accounted for in the relevant review sections (Included studies; Excluded studies; Ongoing studies).

In addition, we searched ClinicalTrials.gov and the WHO International Clinical Trials Registry Platform (ICTRP) for unpublished, planned and ongoing trial reports (21 May 2021), using the search methods detailed in Appendix 1.



Searching other resources

We also searched the reference lists of all retrieved articles.

We did not apply any language restrictions.

Data collection and analysis

For methods used in the previous version of this review, *see* Brown 2002.

For this update, the following methods were used for assessing the reports that were identified as a result of the updated search.

The following methods section of this review is based on a standard template used by Cochrane Pregnancy and Childbirth.

Selection of studies

Two review authors independently assessed for inclusion all the potential studies identified as a result of the search strategy. We resolved any disagreement through discussion or, if required, we consulted the third review author.

Data extraction and management

We designed a form to extract data. For eligible studies, two review authors extracted the data using the agreed form. We resolved discrepancies through discussion and if required, the third review author was consulted. Data were entered into Review Manager 5 software (RevMan 2014) and checked for accuracy.

When information regarding any of the above was unclear, we planned to contact authors of the original reports to provide further details.

Assessment of risk of bias in included studies

Two review authors independently assessed risk of bias for each study using the criteria outlined in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2019). Any disagreement was resolved by discussion or by involving a third assessor.

(1) Random sequence generation (checking for possible selection bias)

For each included study, we described the method used to generate the allocation sequence in sufficient detail to allow an assessment of whether it should produce comparable groups.

We assessed the method as:

- low risk of bias (any truly random process, e.g. random number table; computer random number generator);
- high risk of bias (any non-random process, e.g. odd or even date of birth; hospital or clinic record number);
- unclear risk of bias.

(2) Allocation concealment (checking for possible selection bias)

For each included study, we described the method used to conceal allocation to interventions prior to assignment and assessed whether intervention allocation could have been foreseen in advance of, or during recruitment, or changed after assignment.

We assessed the methods as:

- Cochrane Database of Systematic Reviews
- low risk of bias (e.g. telephone or central randomisation; consecutively numbered sealed opaque envelopes);
- high risk of bias (open random allocation; unsealed or nonopaque envelopes, alternation; date of birth);
- unclear risk of bias.

(3.1) Blinding of participants and personnel (checking for possible performance bias)

For each included study, we described the methods used, if any, to blind study participants and personnel from knowledge of which intervention a participant received. We considered that studies were at low risk of bias if they were blinded, or if we judged that the lack of blinding unlikely to affect results. We assessed blinding separately for different outcomes or classes of outcomes.

We assessed the methods as:

- low, high or unclear risk of bias for participants;
- low, high or unclear risk of bias for personnel.

(3.2) Blinding of outcome assessment (checking for possible detection bias)

For each included study, we described the methods used, if any, to blind outcome assessors from knowledge of which intervention a participant received. We assessed blinding separately for different outcomes or classes of outcomes.

We assessed methods used to blind outcome assessment as:

• low, high or unclear risk of bias.

(4) Incomplete outcome data (checking for possible attrition bias due to the amount, nature and handling of incomplete outcome data)

We described for each included study, and for each outcome or class of outcomes, the completeness of data including attrition and exclusions from the analysis. We stated whether attrition and exclusions were reported and the numbers included in the analysis at each stage (compared with the total randomised participants), reasons for attrition or exclusion where reported, and whether missing data were balanced across groups or were related to outcomes. Where sufficient information was reported, or could be supplied by the trial authors, we planned to re-include missing data in the analyses which we undertook.

We assessed methods as:

- low risk of bias (e.g. no missing outcome data; missing outcome data balanced across groups);
- high risk of bias (e.g. numbers or reasons for missing data imbalanced across groups; 'as treated' analysis done with substantial departure of intervention received from that assigned at randomisation);
- unclear risk of bias.

(5) Selective reporting (checking for reporting bias)

We described for each included study how we investigated the possibility of selective outcome reporting bias and what we found.

We assessed the methods as:



- low risk of bias (where it is clear that all of the study's prespecified outcomes and all expected outcomes of interest to the review have been reported);
- high risk of bias (where not all the study's pre-specified outcomes have been reported; one or more reported primary outcomes were not pre-specified; outcomes of interest are reported incompletely and so cannot be used; study fails to include results of a key outcome that would have been expected to have been reported);
- unclear risk of bias.

(6) Other bias (checking for bias due to problems not covered by (1) to (5) above)

We described for each included study any important concerns we had about other possible sources of bias.

Measures of treatment effect

Dichotomous data

For dichotomous data, we presented results as a summary risk ratio (RR) with its 95% CI.

Continuous data

We presented continuous data using the mean difference (MD) and 95% CIs where the outcome was measured in the same way in different trials. Where different scales were used to measure continuous outcomes, we combined the data using the standardised mean difference (SMD), with the following interpretation:

- small effect: SMD ≥ 0.2 and < 0.5;
- moderate effect: SMD ≥ 0.5 and < 0.8;
- large effect: SMD ≥ 0.8.

Unit of analysis issues

Since all the data come from parallel group RCTs, the unit of analysis for maternal outcomes is per woman randomised. For infant outcomes, the unit of analysis is per infant. For multi-arm trials where there are two groups defined as 'early discharge' and a third defined as 'standard discharge', e.g. with one group discharged at 12 to 24 hours after giving birth, another group discharged at 25 to 48 hours, and a third group discharged at three days, we analysed the two early discharge groups as one combined group.

We did not identify any cluster-RCTs, but if we have data from such trials in future updates, we will include it in the analysis, along with individually randomised trials. We will adjust either their sample sizes or standard errors by using an estimate of the intracluster correlation coefficient (ICC) derived from the trial (if possible), from a similar trial or from a study of a similar population, as described in the *Cochrane Handbook* (Higgins 2019). If we use ICCs from other sources, we will report this and conduct sensitivity analyses to investigate the effect of variation in the ICC. If we identify both cluster-randomised trials and individually randomised trials, we plan to synthesise the relevant information. We will consider it reasonable to combine the results from both if there is little heterogeneity between the study designs and the interaction between the effect of intervention and the choice of randomisation unit is considered to be unlikely.

Dealing with missing data

For included studies, we noted the levels of attrition. Many trials of early postnatal discharge had high levels of missing data due to women being excluded from the study after giving birth, because the circumstances of the birth meant they were no longer eligible for early discharge. We described the extent of missing data in the included studies, clearly distinguishing between data missing due to women no longer meeting the eligibility criteria and data missing due to women who were lost to follow-up, women who chose to withdraw from the trial, or those who were excluded by the trialists for any other reason.

For all outcomes, analyses were carried out, as far as possible, on an intention-to-treat basis, i.e. we attempted to include all participants randomised to each group in the analyses. The denominator for each outcome in each trial was the number randomised, minus any women whose outcomes were known to be missing.

Assessment of heterogeneity

We assessed heterogeneity in each meta-analysis by visual inspection of forest plots and consideration of the I² statistic. Since strict thresholds to interpret I² are not recommended, we used the following as a guide:

- 0% to 40%: might not be important;
- 30% to 60%: may represent moderate heterogeneity;
- 50% to 90%: may represent substantial heterogeneity;
- 75% to 100%: considerable heterogeneity.

Where we identified evidence of heterogeneity, we explored it and described its possible causes.

Assessment of reporting biases

We investigated reporting biases (such as publication bias) using funnel plots for all meta-analyses where 10 or more studies contributed data. When funnel plot asymmetry was detected, it was described and possible causes explored. If asymmetry was suggested by a visual assessment, we performed exploratory analyses to investigate it.

Data synthesis

Statistical analysis was carried out using the Review Manager 5 software (RevMan 2014). We used fixed-effect meta-analysis for combining data where it was reasonable to assume that studies were estimating the same underlying treatment effect: i.e. where trials were examining the same intervention, and the trials' populations and methods were judged sufficiently similar.

If there was clinical heterogeneity sufficient to expect that the underlying treatment effects differed between trials, or if substantial statistical heterogeneity was detected, we used random-effects meta-analysis to produce an overall summary, if an average treatment effect across trials was considered clinically meaningful. The random-effects summary was treated as the average range of possible treatment effects and we discussed the clinical implications of treatment effects differing among trials. Where the average treatment effect was not clinically meaningful, we did not combine trials. If random-effects analyses were used, the

results were presented as the average treatment effect with 95% CIs, and the estimates of Tau² and I².

Subgroup analysis and investigation of heterogeneity

Substantial heterogeneity was investigated using subgroup analyses and sensitivity analyses. Where we identified substantial heterogeneity, we considered whether an overall summary was meaningful, and if it was, we used random-effects analysis.

We carried out the following subgroup analyses.

- Early discharge (less than 24/48 hours) versus later discharge.
- Mode of birth (vaginal versus caesarean section).

To investigate the possible influence of changing clinical practice over time we also carried out post-hoc subgroup analysis according to the decade when the trials were conducted.

Where data were sufficient, we carried out subgroup analyses for the following outcomes.

Infant outcomes

- Infants readmitted for neonatal morbidity (including jaundice, dehydration, infections) within 28 days after birth.
- Infant mortality within a year after birth.

Maternal outcomes

- Women readmitted for complications related to childbirth (including postpartum haemorrhage, retained products of conception, infection, postpartum psychosis) in the first six weeks after giving birth.
- Women scoring above the cut-off score indicating probable depression on a well-validated standardized instrument for measuring depression within six months after giving birth.
- Women breastfeeding (exclusively or partially) at six weeks after giving birth.

We assessed subgroup differences by interaction tests available within RevMan 5 (RevMan 2014) and by investigating the extent of overlap between 95% CIs of the effect estimates belonging to each subgroup. We reported the results of subgroup analyses quoting the P value of the Chi² statistic and the interaction test I² value.

Sensitivity analysis

We performed sensitivity analyses to explore the effect of risk of bias assessed by random sequence generation, allocation concealment, and differential attrition, with studies at high risk of bias in those domains being excluded from the analyses in order to assess whether this made any difference to the overall result.

Sensitivity analyses were also planned to explore the impact of including studies with high levels of missing data, regardless of whether or not the proportion with missing data was balanced across treatment groups. This enabled us to address the issue of women being excluded from analyses after randomisation due to no longer meeting the criteria for early discharge (for example, some studies excluded women after randomisation if they had given birth via caesarean section). We considered a study to have a high level of missing data if 20% or more of the women randomised were not included in the analysis. We performed sensitivity analyses on the following outcomes.

Infant outcomes

- Infants readmitted for neonatal morbidity (including jaundice, dehydration, infections) within 28 days after birth.
- Infant mortality within a year after birth.

Maternal outcomes

- Women readmitted for complications related to childbirth (including postpartum haemorrhage, retained products of conception, infection, postpartum psychosis) in the first six weeks after giving birth.
- Women scoring above the cut-off score indicating probable depression on a well-validated standardized instrument for measuring depression within six months after giving birth.
- Women breastfeeding (exclusively or partially) at six weeks after giving birth.

Summary of findings and assessment of the certainty of the evidence

For this update, we assessed the certainty of the evidence using the GRADE approach, as outlined in the GRADE Handbook (Schünemann 2013). We assessed the certainty of the body of evidence relating to the following outcomes, for the main comparison of a policy of early discharge compared with standard discharge from hospital.

RCT data are initially considered to provide high-certainty evidence but can be downgraded by one or two levels (for serious or very serious concerns, respectively) in the presence of study limitations (high risk of bias), indirectness of evidence, inconsistency of effect estimates and imprecision of effect estimates, as well as the potential of publication bias.

Infant outcomes

- 1. Infants readmitted to hospital for neonatal morbidity (including jaundice, dehydration, infections) within 28 days of birth.
- 2. Infant mortality within one year after birth.
- 3. Number of contacts with health professionals regarding infant health issues within the first four weeks after birth.

Maternal outcomes

- 1. Women readmitted for complications related to childbirth (including postpartum haemorrhage, retained products of conception, infection, postpartum psychosis) in the first six weeks after giving birth.
- 2. Maternal mortality within six weeks after giving birth.
- 3. Women scoring above the cut-off score indicating probable depression on a well-validated standardized instrument for measuring depression within six months after giving birth
- 4. Women breastfeeding (exclusively or partially) at six weeks after giving birth.
- 5. Number of contacts with health professionals regarding maternal health issues within the first six weeks after giving birth.
- 6. Costs of hospital care in the period immediately following the birth up to the time of discharge.



7. Costs of postnatal care following discharge from hospital in the period up to six weeks after the birth, including community midwife, lactation consultant, general practice, specialist and outpatient visits; readmissions to hospital; attendances at day-stay programs; in-patient stays in mother and baby units.

We used the GRADEpro Guideline Development Tool (GRADEpro GDT) to import data from Review Manager 5.3 (RevMan 2014) in order to create 'Summary of findings' tables. A summary of the intervention effect and a measure of certainty for each of the above outcomes was produced using the GRADE approach. The GRADE approach has five considerations (study limitations, consistency of effect, imprecision, indirectness and publication bias) to assess the certainty of the body of evidence for each outcome. The evidence was downgraded from 'high certainty' by one level for serious (or by two levels for very serious) limitations, depending on assessments for risk of bias, indirectness of evidence, inconsistency of effect estimates, imprecision of effect estimates or potential publication bias.

RESULTS

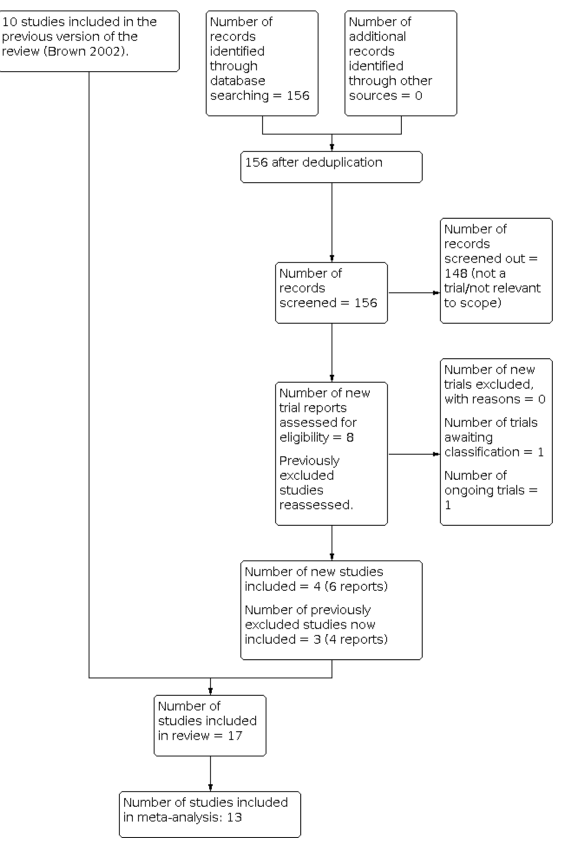
Description of studies

Results of the search

Figure 1 depicts our screening process.



Figure 1. Study flow diagram.





The updated search retrieved 156 reports. We screened out 148 records and assessed eight reports. We also reassessed the excluded studies from the previous version of the review (Brown 2002). We included four new trials (six reports) (Bayoumi 2016; Chiong 2012; Kruse 2021; Taiba 2012), plus three trials that were previously excluded on the basis of lack of relevant outcomes (Burnell 1982; McKeever 2002; Thompson 1999). To adhere to current Cochrane standards we included these three excluded trials in this review because although they did not have useable outcome data, they meet our inclusion criteria.

One trial is ongoing (Namagembe 2014) and we added one trial to Studies awaiting classification for assessment at the next update, as there were no published results available (NCT04422041 2020)

Included studies

We identified 17 trials eligible for inclusion, involving 9409 randomised women. For details of each study, see Characteristics of included studies.

Design

All the included studies were RCTs that randomised women to different treatment groups. We did not identify any cluster-RCTs. One trial was only available in abstract form.

Sample sizes

The largest trial included 3687 women (Bayoumi 2016). The other 16 trials recruited between 50 women (Thompson 1999) and 2257 women (Hellman 1962).

Many trials reported large differences in the numbers of eligible women willing to participate and the numbers of eligible women who declined participation in the trials. Most trials reported the proportion of women eligible to participate in the study in addition to the number of women randomised (Bayoumi 2016; Boulvain 2004; Burnell 1982; Carty 1990; Chiong 2012; Gagnon 1997; Kruse 2021; McKeever 2002; Smith-Hanrahan 1995; Taiba 2012; Thompson 1999; Waldenström 1987). Five of the trials reported a participation rate less than 50% (Boulvain 2004; Gagnon 1997; Kruse 2021; Thompson 1999; Waldenström 1987). A further three trials recruited and randomised between 69% and 90% of women eligible (Burnell 1982; McKeever 2002; Smith-Hanrahan 1995) and the remaining three trials reported 100% uptake to the trial (Carty 1990; Chiong 2012; Taiba 2012).

Setting

The trials included in this review compare 'early postnatal discharge' with standard length of hospital stay as defined in the time and place they were conducted. The trials were undertaken between 1959 (Hellman 1962) and 2019 (Kruse 2021).

We identified no studies from low-income countries. Thirteen studies were from high-income countries; four took place in Canada (Carty 1990; Gagnon 1997; McKeever 2002; Smith-Hanrahan 1995), three in the USA (Brooten 1994; Hellman 1962; Yanover 1976), two in the United Kingdom (Burnell 1982; Winterburn 2000), and one each in Australia (Thompson 1999) Denmark (Kruse 2021), Spain (Sainz Bueno 2005), Sweden (Waldenström 1987), and Switzerland (Boulvain 2004). The remaining three studies took place in middle-income countries; one each in Bangladesh (Taiba 2012), Egypt (Bayoumi 2016) and Malaysia (Chiong 2012).

Participants

The study populations differed considerably among trials. Five trials only included women who were planning, or already had, a caesarean section birth (Bayoumi 2016; Brooten 1994; Chiong 2012; Kruse 2021; Taiba 2012). Ten studies only included women who had given birth vaginally (Burnell 1982; Carty 1990; Gagnon 1997; Hellman 1962; McKeever 2002; Sainz Bueno 2005; Smith-Hanrahan 1995; Thompson 1999; Waldenström 1987; Yanover 1976) and the final two studies included women who had given birth via caesarean section or vaginally (Boulvain 2004; Winterburn 2000).

In nine trials, randomisation took place in pregnancy, usually between 30 weeks and 38 weeks' gestation (Boulvain 2004; Burnell 1982; Carty 1990; Gagnon 1997; Kruse 2021; Thompson 1999; Waldenström 1987; Winterburn 2000; Yanover 1976). Seven trials randomised women immediately after the birth (Bayoumi 2016; Brooten 1994; Chiong 2012; Hellman 1962; McKeever 2002; Sainz Bueno 2005; Smith-Hanrahan 1995). One trial did not describe whether randomisation took place before or after birth.

Most trials specified eligibility criteria designed to limit participation to women at lower risk of complications, although five of the trials included only women who had a caesarean section (Bayoumi 2016; Brooten 1994; Chiong 2012; Kruse 2021; Taiba 2012). In nine trials, pre-randomisation inclusion criteria were designed to select women at low medical risk (Boulvain 2004; Carty 1990; Gagnon 1997; Sainz Bueno 2005; Smith-Hanrahan 1995; Waldenström 1987; Winterburn 2000; Yanover 1976). Eight of these trials recruited both women having their first or subsequent children. The trial conducted by Winterburn and colleagues recruited only women having their first child and planning to breastfeed (Winterburn 2000). Yanover and colleagues specified a number of social eligibility criteria including a requirement that prospective parents currently live together, completion of the final year of high school by mothers, and willingness of fathers to attend prenatal classes (Yanover 1976). Two trials were designed to select women at higher medical risk; the first was the study by Brooten and colleagues that recruited women who had an unplanned caesarean section (Brooten 1994) and the second (Chiong 2012) recruited women after 37 weeks' gestation undergoing planned caesarean section.

Follow-up of participants varied from seven days (Taiba 2012) to one year following childbirth (Bayoumi 2016; Hellman 1962; Waldenström 1987).

Interventions and comparisons

There was substantial heterogeneity in what constituted 'early discharge', to the extent that a length of postnatal hospital stay defined as 'early discharge' in one study was the standard length of postnatal stay in other settings. Standard discharge policies also varied greatly.

We have summarised the interventions and comparisons in Table 1.

Definition of early discharge

The definition of 'early discharge' differed across the trials, reflecting standard practice in the settings in which they were conducted. Three trials defined early as 24 hours or less (Bayoumi 2016; Chiong 2012; Sainz Bueno 2005), one as 28 hours or less (Kruse 2021) and one as 36 hours or less (Thompson 1999). Five trials defined early discharge as a range: between six hours



and 36 hours (Gagnon 1997); between six hours and 48 hours (Winterburn 2000); between 12 hours and 48 hours (Yanover 1976); between 24 and 36 hours (McKeever 2002); and between 24 and 48 hours (Waldenström 1987).

One trial defined early discharge as between 24 and 48 hours for vaginal birth and between 72 and 84 hours after caesarean section (Boulvain 2004).

One trial allocated women to two different early discharge groups; one for discharge at 12 to 24 hours and another for discharge at 25 to 48 hours (Carty 1990).

Five of the 17 trials defined early discharge as 48 hours or longer after giving birth (Brooten 1994; Burnell 1982; Hellman 1962; Smith-Hanrahan 1995; Taiba 2012).

Definition of standard care

Most trials included in the review stated a standard postnatal length of stay at least 48 hours after birth (Bayoumi 2016; Boulvain 2004; Brooten 1994; Burnell 1982; Carty 1990; Chiong 2012; Gagnon 1997; Hellman 1962; Kruse 2021; Sainz Bueno 2005; Smith-Hanrahan 1995; Taiba 2012; Waldenström 1987; Winterburn 2000; Yanover 1976). One study did not state the standard length of stay for the control group (McKeever 2002) and the final trial reported a standard care length of stay as between 37 hours and 48 hours after birth (Thompson 1999).

In several studies, the duration of hospital stay in the comparator group was equal to or shorter than the definition of early discharge in other studies. For instance, women in the comparator groups in four vaginal birth-only studies (Gagnon 1997; Sainz Bueno 2005; Smith-Hanrahan 1995; Thompson 1999) were allocated to hospital stays that were shorter than or equal to the early discharge period of up to 72 hours in one other vaginal birth-only study (Hellman 1962). Similarly, in two caesarean-only studies (Bayoumi 2016; Kruse 2021) early discharge was shorter than standard discharge of 72 hours in another caesarean-only study (Taiba 2012).

Co-interventions

The intensity of antenatal preparation and frequency of midwife home care following discharge offered to women in intervention and control groups differed considerably among trials. Two studies included antenatal home visits for women randomised to early discharge (Carty 1990; Waldenström 1987), and a third offered a prenatal 'preparation for discharge' class for women in both intervention and control arms of the study (Yanover 1976). Home visits by study nurses or nurse-midwives were made to women in the early discharge arms of 13 trials, although in the largest trial (Hellman 1962) study nurses were intended to collect information only and were requested not to provide actual nursing care or support. There was substantial variation in the nature and extent of nurse-midwife support specified in study protocols. Some trials offered a mixture of home visits and phone calls (Boulvain 2004; Gagnon 1997; Smith-Hanrahan 1995) while others included a home visit during pregnancy as well as home visits after the birth (Carty 1990; Waldenström 1987). Six trials restricted midwife home visits to the early discharge group (Brooten 1994; Sainz Bueno 2005; Smith-Hanrahan 1995; Waldenström 1987; Winterburn 2000; Yanover 1976). The other four trials provided a limited number of midwife home visits to women in the control group (Boulvain 2004; Carty 1990; Hellman 1962) or provided home visits on referral by a physician (Gagnon 1997). Two studies did not provide home visits to either arm, but participants could access the clinics and acute care facilities as required (Bayoumi 2016; Chiong 2012). None of the studies provide detailed information about access to primary care services in the settings in which the studies were conducted.

Outcomes

Thirteen trials reported our primary outcomes:

- Infants readmitted for neonatal morbidity (Bayoumi 2016; Boulvain 2004; Brooten 1994; Chiong 2012; Hellman 1962; Kruse 2021; Sainz Bueno 2005; Smith-Hanrahan 1995; Waldenström 1987; Yanover 1976).
- Women readmitted for complications related to childbirth in the first six weeks after giving birth (Bayoumi 2016; Boulvain 2004; Carty 1990; Chiong 2012; Hellman 1962; Kruse 2021; Sainz Bueno 2005; Waldenström 1987; Yanover 1976).
- Women scoring above the cut-off score, indicating probable depression on a well-validated standardized instrument for measuring depression within six months after giving birth (Bayoumi 2016; Boulvain 2004; Chiong 2012; Sainz Bueno 2005; Waldenström 1987).
- Women breastfeeding at six weeks, 12 weeks and six months after giving birth (Bayoumi 2016; Boulvain 2004; Carty 1990; Chiong 2012; Gagnon 1997; Hellman 1962; Sainz Bueno 2005; Waldenström 1987; Winterburn 2000).

The following outcomes were not reported by any of the studies.

- Infants attending hospital casualty or emergency department within seven days, and the first 28 days after the birth.
- Number of contacts with health professionals regarding infant health issues within seven days after the birth.
- Maternal mortality within six weeks.
- Duration of readmissions for women readmitted within the first six weeks after the birth.
- Total duration of maternal hospitalisation over the first six weeks after the birth.
- Women attending hospital casualty or emergency department within the first six weeks after the birth.

Funding sources

Ten trials reported that their funding was from government, university or hospital sources (Boulvain 2004; Brooten 1994; Chiong 2012; Gagnon 1997; Hellman 1962; McKeever 2002; Smith-Hanrahan 1995; Waldenström 1987; Winterburn 2000; Yanover 1976).

Five trials did not report any information about their funding sources (Bayoumi 2016; Burnell 1982; Carty 1990; Sainz Bueno 2005; Taiba 2012).

Two trials reported funding from other sources: Brodene Hartmanns Foundation (Kruse 2021); the Salaried Specialists (Thompson 1999).

Declarations of interest

Fourteen trials did not report any information about the authors' declarations of interest (Boulvain 2004; Brooten 1994; Burnell 1982;



Carty 1990; Gagnon 1997; Hellman 1962; McKeever 2002; Sainz Bueno 2005; Smith-Hanrahan 1995; Taiba 2012; Thompson 1999; Waldenström 1987; Winterburn 2000; Yanover 1976).

Three trials explicitly stated that the authors had no conflicts of interest or had nothing to declare (Bayoumi 2016; Chiong 2012; Kruse 2021).

No trials reported any conflicts of interest.

Missing data

Six trials reported the numbers of women excluded from analysis because they no longer met either the trial eligibility criteria or the criteria for early discharge. The proportion of women excluded for this reason ranged from 0.22% (Boulvain 2004) to 44% (Gagnon 1997). The number of women excluded for this reason was generally balanced between the treatment groups in the trials, with the exception of one trial (Thompson 1999) where 13/24 and 5/24 women were excluded because they no longer fulfilled the early discharge criteria. Eight trials did not report whether any women were excluded because they no longer met either the trial eligibility criteria or the criteria for early discharge (Bayoumi 2016; Brooten 1994; Burnell 1982; Carty 1990; Hellman 1962; Sainz Bueno 2005; Taiba 2012; Yanover 1976). Three trials reported that no women were excluded for no longer meeting either the trial eligibility criteria or the criteria for early discharge (Kruse 2021; McKeever 2002; Winterburn 2000).

All but four trials (Brooten 1994; Burnell 1982; Hellman 1962; Taiba 2012) reported the number of women who did not adhere to the treatment group to which they were allocated, i.e. the number of women who either stayed beyond the early discharge time or who were discharged before the standard discharge time. The proportion of women not adhering to the treatment group ranged from zero (McKeever 2002; Yanover 1976) to 61% (Boulvain

2004). The number of women not adhering to the treatment group was generally balanced between the treatment groups in the trials, except in one trial (Winterburn 2000), where 90/121 and 20/127 women either stayed for a longer or a shorter time than the treatment to which they were allocated. Six trials reported that the women who did not adhere to their treatment group were included in the final analysis according to their allocated group (Boulvain 2004; Chiong 2012; Gagnon 1997; Kruse 2021; Sainz Bueno 2005; Winterburn 2000). One trial (Waldenström 1987) analysed all women in the control group according their allocated group, regardless of whether or not they adhered to it; but this trial analysed the women in the early discharge on a per-protocol basis, i.e. those who did not adhere were not included in the analysis.

Nine trials had a high level of missing data, where 20% or more the women who were randomised were not included in the final analysis (Bayoumi 2016; Carty 1990; Gagnon 1997; McKeever 2002; Smith-Hanrahan 1995; Taiba 2012; Thompson 1999; Waldenström 1987; Yanover 1976). Five trials had no missing data (Brooten 1994; Burnell 1982; Hellman 1962; Kruse 2021; Sainz Bueno 2005), although one of these did not report the number of women who were originally randomised; therefore, it is not clear if any data were missing from the final analysis (Brooten 1994).

Excluded studies

We excluded three trials because they were not designed to measure the effect of early discharge compared with longer length of hospital stay (see Characteristics of excluded studies).

Risk of bias in included studies

The summary risk of bias across all studies for each domain is presented in Figure 2. Risk of bias per domain, per study is presented in Figure 3.

Figure 2. Risk of bias graph: review authors' judgements about each risk of bias item presented as percentages across all included studies.

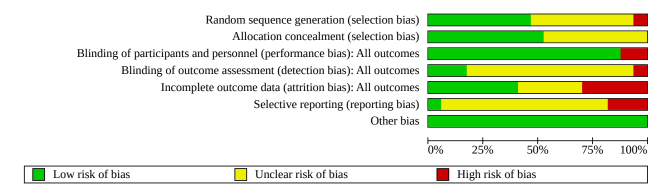




Figure 3. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.

Bayoumi 2016 ? 1 1 2 ? 1 Boulvain 2004 1 1 1 1 ? 1 1 Brooten 1994 1 1 1 1 ? 1 1 ? 1 1 Burnell 1982 ? 2 1 ? 1 ? 1 1 ? 1 1 ? 1 1 ? 1 1 ? 1 1 ? 1 ? 1 ? 1 ? 1 ? 1 ? 1 ?		Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding of participants and personnel (performance bias): All outcomes	Blinding of outcome assessment (detection bias): All outcomes	Incomplete outcome data (attrition bias): All outcomes	Selective reporting (reporting bias)	Other bias
Brooten 1994 Image: Constraint of the sector of the se	-	-		-	_			+
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Kruse 2021 +	-					?		
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Smith-Hanrahan 1995 ?		+		+	+	•	•	+
Taiba 2012 ?	Sainz Bueno 2005	?	+	+	?	?	?	+
Thompson 1999 + <	Smith-Hanrahan 1995	?	?	+	?	+	?	+
Waldenström 1987 ?	Taiba 2012	?	?	+	?	•	?	+
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	Waldenström 1987				-		-	_
		?		+	?	•		Ŧ
Yanover 1976 <mark>? Ŧ ? ?</mark> Ŧ	Winterburn 2000	?	?		?	•		+

See Characteristics of included studies for full details.



Allocation

Random sequence generation

Random sequence generation was detailed in six of the trials in sufficient detail to be judged at low risk of bias (Boulvain 2004; Brooten 1994; Carty 1990; Chiong 2012; Gagnon 1997; Kruse 2021; McKeever 2002; Thompson 1999). Random sequence generation was not clear in eight of the trials (Bayoumi 2016; Burnell 1982; Sainz Bueno 2005; Smith-Hanrahan 1995; Taiba 2012; Waldenström 1987; Winterburn 2000; Yanover 1976). We judged one trial as high risk of bias because it did not use any randomisation process to allocate women to the intervention group, while the control group allocation was done in a randomised fashion that was not described in detail (Hellman 1962).

Allocation concealment

We judged nine trials as low risk of bias because methods to ensure allocation concealment were clearly documented (Bayoumi 2016; Boulvain 2004; Brooten 1994; Carty 1990; Chiong 2012; Gagnon 1997; Kruse 2021; Sainz Bueno 2005; Thompson 1999). There was insufficient information available to determine the adequacy of concealment prior to randomisation for the other eight trials so we judged them as unclear risk of bias (Burnell 1982; Hellman 1962; McKeever 2002; Smith-Hanrahan 1995; Taiba 2012; Waldenström 1987; Winterburn 2000; Yanover 1976).

Blinding

Blinding of participants and caregivers

It was not possible to blind participants or caregivers to allocation status for this intervention. However, studies were judged at low risk of bias if the main outcomes were unlikely to be affected by lack of blinding (for example, readmission rates to hospital) and high risk if the main outcomes were likely to be affected by the lack of participant blinding (for example, participant satisfaction). Fifteen studies were considered to be at low risk of performance bias (Bayoumi 2016; Boulvain 2004; Brooten 1994; Carty 1990; Chiong 2012; Gagnon 1997; Hellman 1962; Kruse 2021; McKeever 2002; Sainz Bueno 2005; Smith-Hanrahan 1995; Taiba 2012; Waldenström 1987; Winterburn 2000; Yanover 1976) and a further two studies were judged at high risk of performance bias due to lack of participant blinding (Burnell 1982; Thompson 1999).

Blinding of outcome assessors

We judged three studies as low risk of detection bias because clear methods to blind the outcome assessment to participant allocation were described (Bayoumi 2016; McKeever 2002; Thompson 1999). It was unclear whether outcome assessment was blinded to allocation status in a further 13 studies (Boulvain 2004; Brooten 1994; Burnell 1982; Carty 1990; Chiong 2012; Gagnon 1997; Hellman 1962; Sainz Bueno 2005; Smith-Hanrahan 1995; Taiba 2012; Waldenström 1987; Winterburn 2000; Yanover 1976). We judged one study as high risk of detection bias because the outcome assessors were not blinded to participant allocation (Kruse 2021).

Incomplete outcome data

Low attrition rates, which were balanced across both arms of the trial, were described in seven of the studies and we judged them to be at low risk of bias (Bayoumi 2016; Boulvain 2004; Chiong 2012; Kruse 2021; Smith-Hanrahan 1995; Thompson 1999; Winterburn 2000). Levels of attrition and how missing data were accounted for was not clear in five trials and we judged these as unclear risk of bias (Brooten 1994; Burnell 1982; Hellman 1962; Sainz Bueno 2005; Yanover 1976) The remaining six trials we judged at high risk of attrition bias either due to differential attrition rates between the early discharge group and standard care group (Carty 1990; Gagnon 1997; McKeever 2002; Taiba 2012; Waldenström 1987) or high attrition rates in the trial overall (Smith-Hanrahan 1995).

Selective reporting

None of the studies had published protocols, although the majority of included studies were published at a time when this was not a requirement. We judged these studies as unclear risk of reporting bias (Bayoumi 2016; Boulvain 2004; Brooten 1994; Carty 1990; Chiong 2012; Gagnon 1997; Hellman 1962; Sainz Bueno 2005; Smith-Hanrahan 1995; Taiba 2012; Waldenström 1987; Winterburn 2000; Yanover 1976). Two of the trials did not report outcomes that were specified in the methods section and we judged these to be at high risk of reporting bias (Burnell 1982; McKeever 2002; Thompson 1999). Kruse 2021 and colleagues did not publish a protocol but the trial was registered prospectively and not all pre-specified outcomes were reported in full; however, correspondence with the trial author confirmed that the outcomes were measured and the data analysis is in progress. We judged this trial to be at low risk of reporting bias.

Other potential sources of bias

All 17 studies included in the review appeared to be at low risk of any other sources of bias.

Effects of interventions

See: Summary of findings 1 Early discharge compared to standard discharge for healthy mothers and term infants

Primary infant outcomes

Infants readmitted for neonatal morbidity (including jaundice, dehydration, infections) within seven days

We are uncertain if early discharge, compared with standard discharge, has any effect on the risk of infant readmission within seven days (RR 1.15, 95% CI 0.42 to 3.16; 247 participants; two studies; $I^2 = 0\%$; Analysis 1.1).

Infants readmitted for neonatal morbidity (including jaundice, dehydration, infections) within 28 days

Early discharge probably slightly increases the number of infants readmitted within 28 days compared to infants in the standard care group (RR 1.59, 95% Cl 1.27 to 1.98; 6918 participants; 10 studies; $l^2 = 12\%$; Analysis 1.2; moderate-certainty evidence; Summary of findings 1). The risk of infant readmission in the early discharge group was 69 per 1000 infants compared to 43 per 1000 infants in the standard care group.

The tests for subgroup differences did not suggest evidence of a difference in effect between different decades when the trials took place (P = 0.24, l^2 = 26.3%), different modes of birth (P = 0.65, l^2 = 0%; Analysis 1.3) or the varying definitions of early discharge (P = 0.55, l^2 = 0%; Analysis 1.4).

The sensitivity analysis, removing two trials at high risk of bias in the random sequence generation and differential attrition



domains (Hellman 1962; Waldenström 1987), did not change the effect estimate substantially (RR 1.58, 95% Cl 1.26 to 1.99; 4663 participants; $l^2 = 33\%$).

A second sensitivity analysis, removing trials with a high level of missing data (defined as 20% or more of the infants of women randomised were not included in the analysis), suggests that it is uncertain whether early discharge, compared with standard discharge, has any effect on the risk of infant readmission within 28 days (RR 1.10, 95% CI 0.67 to 1.81; 3647 participants; $I^2 = 16\%$). Four trials were removed in order to conduct this sensitivity analysis (Bayoumi 2016; Smith-Hanrahan 1995; Waldenström 1987; Yanover 1976).

Primary maternal outcomes

Women readmitted for complications related to childbirth (including postpartum haemorrhage, retained products of conception, infection, postpartum psychosis) within six weeks postpartum

Early discharge probably results in little to no difference in women readmitted within six weeks but the 95% CI suggests that the true effect may be either an increase or decrease in risk (RR 1.12, 95% CI 0.82 to 1.54; 6992 participants; 11 studies; $I^2 = 0\%$; moderatecertainty evidence; Analysis 1.5; Summary of findings 1).

The tests for subgroup differences did not suggest evidence of a difference in effect between different decades when the trials took place (P = 0.25, $I^2 = 25.4\%$), mode of birth (P = 0.69; $I^2 = 0\%$; Analysis 1.6) or definition of early discharge (P = 0.66; $I^2 = 0\%$; Analysis 1.7).

Sensitivity analysis removing three trials at high risk of bias in the random sequence generation, allocation concealment or differential attrition domains (Carty 1990; Hellman 1962; Waldenström 1987), did not change the effect estimate substantially (RR 1.07, 95% CI 0.77 to 1.48; 4663 participants; $I^2 =$ 0%).

A second sensitivity analysis, removing three trials with a high level of missing data (defined as 20% or more of the women randomised were not included in the analysis) (Bayoumi 2016; Carty 1990; Waldenström 1987) made the effect estimate more imprecise, with a substantially wider 95% CI (RR 1.28, 95% CI 0.69 to 2.40; 3590 participants; $I^2 = 0\%$).

Women scoring above the cut-off score indicating probable depression on a well-validated standardized instrument for measuring depression within six months after the birth

Early discharge may result in little to no difference in the risk of depression within six months but the wide 95% CI suggests the possibility that the true effect is either an increase or a reduction in

risk (RR 0.80, 95% CI 0.46 to 1.42; 4333 participants; five studies; I² = 58%; Analysis 1.8; low-certainty evidence; Summary of findings 1).

The tests for subgroup differences did not suggest evidence of a difference in effect between different decades when the trials took place (P = 0.46, $I^2 = 0\%$), different modes of birth (P = 0.41, $I^2 = 0\%$; Analysis 1.9) or time of early discharge (P = 0.79, $I^2 = 0\%$, Analysis 1.10).

Sensitivity analysis removing one trial at high risk of bias in the differential attrition domain (Waldenström 1987) did not change the effect estimate substantially (RR 0.81, 95% CI 0.43 to 1.53).

A second sensitivity analysis removing trials with a high level of missing data, where 20% or more of the women randomised were not included in the analysis (Bayoumi 2016; Waldenström 1987), did not change the effect estimate substantially (RR 0.62, 95% CI 0.34 to 1.12).

Additionally, one study reported mean scores on the Beck Depression Inventory at one month postpartum favouring early discharge (Carty 1990). The mean score for women randomised to discharge within 24 hours of the birth was 4.5 (standard deviation (SD) 2.54) versus a mean score of 7.8 (SD 6.46) for women randomised to discharge four or more days after the birth (P < 0.05). Higher scores on this scale indicate poorer emotional well-being. It was not possible to include these data in the meta-analysis because there was insufficient data provided in the paper.

Women breastfeeding (exclusively or partially) at six weeks after giving birth

Early discharge probably results in little to no difference in women breastfeeding at six weeks after giving birth (RR 1.04, 95% CI 0.96 to 1.13; 7156 participants; 10 studies; $I^2 = 79\%$; Analysis 1.11; moderate-certainty evidence; Summary of findings 1).

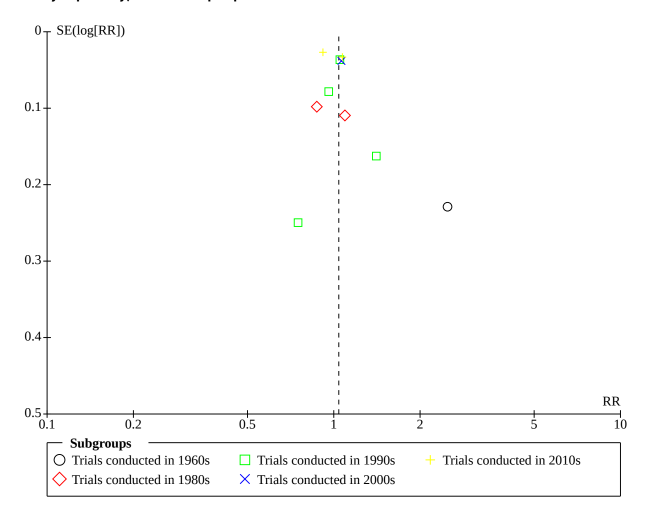
The test for subgroup differences suggested a difference in effect between trials taking place in different decades (P = 0.004, l^2 = 73.8%). This difference appears to be due to one trial that took place in the early 1960s where more women in the early discharge group were breastfeeding at six weeks compared to women in the standard discharge group, while the trials taking place in the decades since the 1960s did not observe a clear difference between groups.

The tests for subgroup differences did not suggest evidence of a difference in effect between different modes of birth (P = 0.61, I² = 0%, Analysis 1.12) or time of early discharge did not indicate evidence of a difference in effect between groups (P = 0.74, I² = 0%, overlapping Cls; Analysis 1.13).

The symmetry of the funnel plot (Figure 4) did not suggest evidence of data missing due to non-publication of studies with statistically insignificant findings.



Figure 4. Funnel plot of comparison: 1 Early versus standard discharge, outcome: 1.13 Women breastfeeding (exclusively or partially) at six weeks postpartum.



Sensitivity analysis removing four trials at high risk of bias in the random sequence generation, allocation concealment or differential attrition domains (Carty 1990; Chiong 2012; Hellman 1962; Waldenström 1987) did not substantially change the effect estimate (RR 1.01, 95% CI 0.92 to 1.09).

A second sensitivity analysis removing five trials with a high level of missing data (defined as 20% or more of the women randomised were not included in the analysis) (Bayoumi 2016; Carty 1990; Gagnon 1997; Smith-Hanrahan 1995; Waldenström 1987), did not change the effect estimate substantially (RR 1.09, 95% CI 0.98 to 1.21).

Women breastfeeding (exclusively or partially) 12 weeks after giving birth

One study measured the number of women breastfeeding at 12 weeks after giving birth and found more women in the early discharge group were breastfeeding (RR 1.21, 95% CI 1.03 to 1.41; 430 participants; Analysis 1.14).

Women breastfeeding (exclusively or partially) at six months after giving birth

There may be little to no difference between early discharge and later discharge in the number of women breastfeeding at six months (RR 1.12, 95% Cl 0.94 to 1.32; 973 participants; three studies; $l^2 = 52\%$; Analysis 1.15). To investigate the moderate level of heterogeneity in the analysis we performed the analysis again with random-effects and the effect estimate did not change substantially (RR 1.11, 95% Cl 0.87 to 1.43).

Secondary infant outcomes

Infant mortality within 28 days

It is uncertain whether early discharge, compared with standard discharge, has any effect on the risk of infant mortality within 28 days (RR 0.39, 95% CI 0.04 to 3.74; 4882 participants; two studies; $I^2 = 0\%$; Analysis 1.16).

Infant mortality within one year

It is uncertain whether early discharge, compared with standard discharge, has any effect on the risk of infant mortality within one year because the certainty of evidence is low and 95% CI is

consistent with possible benefit and possible harm (RR 0.45, 95% CI 0.07 to 2.77; 4984 participants; three studies; I² = 0%; Analysis 1.17; low-certainty evidence; Summary of findings 1).

Subgroup analysis on the basis of mode of birth was not possible because there was only trial in the caesarean birth group and it had zero events in both arms. For the same reason we could not perform subgroup analysis with regard to time of early discharge.

Sensitivity analysis removing one trial at high risk of bias due to random sequence generation (Hellman 1962) did not change the effect estimate substantially but made the CI wider (RR 0.36, 95% CI 0.02 to 8.63). Similarly, sensitivity analysis removing one trial with high levels of missing data (Waldenström 1987) did not change the effect estimate substantially but made the CI wider (RR 0.52, 95% CI 0.06 to 4.64).

Duration of infant readmissions for infants readmitted within seven days after the birth

One study reported that the single infant in the study that was readmitted within seven days stayed in hospital for two days (Waldenström 1987). The infant was in the early discharge group (n = 50). No infants in the later discharge group were readmitted (n = 54).

No other studies reported duration of infant readmissions in the first seven days.

Duration of infant readmissions for infants readmitted within the first 28 days after the birth

One study reported that 1/50 and 1/54 infants each spent two days in hospital in the first 28 days (Waldenström 1987).

No other studies reported duration of infant readmissions in the first 28 days.

Total duration of infant hospitalisation over the first 28 days

One study reported total duration of infant hospitalisation over the first 28 days. However it only reported the mean number of days for the control group (two days). No data were reported for mean duration of infant hospitalisation for the early discharge group (Waldenström 1987).

Infants attending hospital casualty or emergency department within seven days, and the first 28 days after the birth

Not reported.

Number of contacts with health professionals regarding infant health issues within seven days after the birth

Not reported.

Number of contacts with health professionals regarding infant health issues within the first four weeks after the birth

Early postnatal discharge, compared with later discharge, probably makes little to no difference in the number of infants having at least one unscheduled medical consultation or contact with health professionals within the first four weeks after birth (RR 0.88, 95% CI 0.67 to 1.16; 639 participants; three studies; I² = 0%; Analysis 1.18; moderate-certainty evidence; Summary of findings 1) (Brooten 1994; Chiong 2012; Gagnon 1997).

Two other studies reported data that could not be included in the meta-analysis (Kruse 2021; Waldenström 1987). The results indicated little to no difference between the groups in terms of number of contacts with health professionals regarding infant health issues (Table 2).

Secondary maternal outcomes

Maternal mortality within six weeks postpartum

Not reported.

Maternal mortality within one year postpartum

Two studies reported maternal mortality within one year (Bayoumi 2016; Taiba 2012). Of 1545 women allocated to early discharge and 1653 women allocated to standard discharge, there were no maternal deaths within one year.

Duration of readmissions for women readmitted within the first six weeks after the birth

Not reported.

Total duration of maternal hospitalisation over the first six weeks after the birth

Not reported.

Women attending hospital casualty or emergency department within first six weeks after the birth

Not reported.

Number of contacts with health professionals regarding maternal health issues within the first six weeks after the birth

Two studies reported the number of women having at least one unscheduled medical consultation or contact with health professional (Brooten 1994; Chiong 2012), with both studies including women following caesarean section. There is probably little to no difference in the number of women who have at least one unscheduled medical consultation after having been discharged early, compared with later discharge (RR 0.72, 95% CI 0.43 to 1.20; 464 participants; two studies; I² = 29%; Analysis 1.19; moderatecertainty evidence; Summary of findings 1).

Two other studies reported data suggesting little difference between the groups in terms of contact with health professionals for maternal health issues but they were not suitable for inclusion in the meta-analysis (Boulvain 2004; Kruse 2021; see Table 3).

Women reporting tiredness or exhaustion in the first six weeks after the birth

Three trials reported measures of tiredness and exhaustion in the first six weeks after the birth but they reported no data that were suitable for pooled analysis. Waldenström and colleagues compared mean values on a five-point scale that rated fatigue and alertness from 'very alert' to 'very tired' for the first 14 days postpartum. No evidence of a difference was found between groups, but women in both groups were most tired on the day following discharge (Waldenström 1987). Smith-Hanrahan and colleagues used a 10 centimetre visual analogue scale with anchors ranging from zero (not tired, full of energy, peppy) to 10 (total exhaustion) with assessments at two to three days, one week and six weeks after the birth. No significant differences in



mean scores were found between intervention and control groups (Smith-Hanrahan 1995). Sainz Bueno and colleagues report the increase in puerperal fatigue at one month compared with one week postpartum and found no differences between groups (Sainz Bueno 2005).

Women reporting physical health problems (including perineal pain, perineal infection, breast soreness, breast infection, caesarean wound pain, caesarean wound infection) in the first six weeks after the birth

In one trial fewer women in the early discharge group reported physical health problems in the first six weeks after giving birth compared with the standard discharge group (RR 0.25, 95% CI 0.11 to 0.59; 200 participants; one study; Analysis 1.20; Taiba 2012).

Women reporting that they lacked confidence about caring for their baby in the first month and the first six months after being discharged from hospital

Not reported.

Women reporting infant feeding problems in the first four weeks after the birth

Two trials reported data on infant feeding problems (Boulvain 2004; Hellman 1962). We are uncertain whether there is any difference between early discharge and standard care for women reporting infant feeding problems in the first four weeks after the birth (RR 0.89, 95% CI 0.43 to 1.86; 2405 participants; two studies; I² = 88%; Analysis 1.21). Random effects analysis was used to combine trials, due to substantial heterogeneity, which may be due to different modes of births in different trial populations. The estimate of fixedeffect analysis was not substantially different: RR 0.87, 95% CI 0.68 to 1.11).

Women reporting they received conflicting advice regarding breastfeeding in the first six weeks after birth

Not reported.

Women who express satisfaction with their postnatal care (including in hospital and/or post-discharge care)

Six trials used various methods to measure women's satisfaction with postnatal care in hospital (Boulvain 2004; Carty 1990; Gagnon 1997; Hellman 1962; Sainz Bueno 2005; Waldenström 1987). Four studies used dichotomous measures (Boulvain 2004; Hellman 1962; Sainz Bueno 2005; Waldenström 1987) and two used numerical scales (Carty 1990; Gagnon 1997).

There may be little to no difference in number of women satisfied between women who were discharged early and women discharged according to standard protocol (RR 1.10, 95% CI 0.95 to 1.29; 3098 participants; four studies; $I^2 = 93\%$; Analysis 1.22).

The high I² statistic may be due to variations in the trials' definition of early discharge and/or standard discharge. The early discharge groups ranged from discharge within 24 hours (Sainz Bueno 2005), between 24 and 48 hours (Boulvain 2004; Waldenström 1987), to up to 72 hours (Hellman 1962). In all of the studies standard discharge was considerably later than early discharge; two to three days later (Boulvain 2004), one day later (Sainz Bueno 2005), at least two days later (Hellman 1962), four to five days later (Waldenström 1987). One study (Carty 1990) measured women's satisfaction with nursing care using a questionnaire with a maximum score of 110 (higher score = greater satisfaction). Another study (Gagnon 1997) used a 5-point Likert scale to measure women's satisfaction with health care in first day 10 days postpartum (higher score = greater satisfaction). Combining these studies using standardized mean difference indicated substantially greater satisfaction with early discharge compared with standard discharge (SMD 0.74, 95% CI 0.50 to 0.98; 306 participants; two studies; $I^2 = 0\%$; Analysis 1.23).

No trials included measures specifically focusing on women's views of postnatal care after they left hospital.

Women who perceive their length of hospital stay as too short

Only one trial reported on the proportion of women who perceived their length of hospital as too short (Yanover 1976). Very few women in the trial conducted by Yanover and colleagues thought their length of stay too short (2/41 in the early discharge group and 1/41 in the group randomised to standard care (RR 2.00, 95% CI 0.19 to 21.21; 82 participants; Analysis 1.24).

Women who perceive their length of hospital stay as too long

Two trials reported on the proportion of women who perceived their length of stay as too long. Yanover and colleagues found slightly fewer women in the early discharge group that thought their stay was 'too long' (5/41 in the early discharge group versus 9/41 in the standard care group (RR 0.56, 95% CI 0.20 to 1.52; 82 participants; Analysis 1.25). One study (Sainz Bueno 2005) only reported the proportion of women who perceived their length of stay as too long in the control group (54/125) but no data were reported for the intervention group.

Partner outcomes

None of the trials reported any of our pre-specified partner outcomes. One trial measured outcomes related to fathers' involvement in infant care but it did not report any data that we could use for analysis (Waldenström 1987). They found that fathers' involvement was higher in the first few days after the birth in the early discharge group, both among first-time fathers and those having their second or a subsequent child. First-time fathers in the early discharge group spent an average of 183 minutes per day with their baby in the first two to four days, compared with 71 minutes for the control group (student's t test = 4.5, P < 0.001). Fathers with two or more children spent an average of 89 minutes with the new baby in the early discharge group, compared with 44 minutes in the control group (t = 2.9, P < 0.01). These differences were no longer apparent at two weeks and six weeks after the birth.

Economic outcomes

Costs of hospital care in the period immediately following the birth up to the time of discharge

Three trials reported mean costs of hospitalisation (see Table 4; Boulvain 2004; Brooten 1994; Sainz Bueno 2005). Early discharge may slightly reduce the costs of hospital care in the period immediately following the birth up to the time of discharge (lowcertainty evidence; Summary of findings 1).

Costs of postnatal care following discharge from hospital in the period up to six weeks after the birth, including community midwife, lactation consultant, general practice, specialist and

outpatient visits; readmissions to hospital; attendances at daystay programs; in-patient stays in mother and baby units

Three trials reported data on costs of postnatal care (see Table 5; Boulvain 2004; Brooten 1994; Sainz Bueno 2005). Early discharge may result in little to no difference in costs of postnatal care following discharge from hospital in the six weeks after the birth (low-certainty evidence; Summary of findings 1).

One trial reported that early discharge was associated with a higher mean cost of all postnatal care and a higher mean cost of community care but that the overall mean cost of all care, including hospitalisation, was lower with early discharge than later discharge (Boulvain 2004). The same trial also measured non-medical costs attributable to health care in the first six weeks after birth (travel to healthcare providers, childcare support for siblings) and also the costs associated with loss of income if a partner required time off work. They found no differences between the groups in these non-medical and indirect costs.

Costs of practical support following discharge from hospital in the period up to six weeks after the birth, including paid and unpaid home help, care of the baby and of siblings

Not reported.

DISCUSSION

Summary of main results

The 17 trials we identified, with 9409 randomised women, help to consolidate the evidence base on the safety of early postnatal discharge for women and infants.

Infant outcomes

Early discharge of healthy mothers and term infants probably slightly increases the number of infants readmitted within 28 days for neonatal morbidity (moderate-certainty evidence). It is uncertain whether early discharge has any effect on the risk of infant mortality within 28 days (low-certainty evidence). Early postnatal discharge probably makes little to no difference in the number of infants having at least one unscheduled medical consultation or contact with health professionals within the first four weeks after birth.

Maternal outcomes

Early discharge probably results in little to no difference in women readmitted within six weeks postpartum for complications related to childbirth (moderate-certainty evidence) but the wide 95% CI indicates the possibility that the true effect is either an increase or decrease in risk. Similarly, early discharge may result in little to no difference in the risk of depression within six months postpartum (low-certainty evidence) but the wide 95% CI suggests the possibility that the true effect is either an increase or a reduction in risk.

Early discharge probably results in little to no difference in women breastfeeding at six weeks postpartum (moderatecertainty evidence) or in the number of women having at least one unscheduled medical consultation or contact with health professionals (moderate-certainty evidence).

Maternal mortality within six weeks postpartum was not reported in any of the studies.

Costs

Early discharge may slightly reduce the costs of hospital care in the period immediately following the birth up to the time of discharge (low-certainty evidence) but it may result in little to no difference in costs of postnatal care following discharge from hospital in the period up to six weeks after the birth (low-certainty evidence).

Overall completeness and applicability of evidence

We identified evidence for all but one of our pre-specified GRADE outcomes; none of the studies reported maternal mortality within six weeks postpartum. It is not clear if this is because the outcome was not measured or if there were no deaths and the trialists simply have not reported such a rare outcome in the settings where the trials were conducted.

Several of our pre-specified secondary outcomes were not reported at all, and there is a notable absence of evidence relating to partner outcomes. Furthermore, it is difficult to draw conclusions and generalise from the limited costs data that we identified.

The definition of early discharge differed considerably between trials and mirrored the differences in standard discharge policies in each context. In most of the trials included in this review, early discharge was accompanied by some level of post-discharge nursing or midwifery support. In practice, policies promoting shorter length of stay may not always be implemented with accompanying primary care support in the days following discharge. It remains unclear how important home midwifery or nursing support is to the safety and acceptability of early discharge programs.

Quality of the evidence

We assessed the risk of bias in terms of random sequence generation to be generally low or unclear. Risk of bias related to blinding of participants was generally low because we judged that lack of blinding would be unlikely to affect the outcomes measured. In contrast, risk of bias related to outcome assessment was unclear because most trials did not report whether or not outcome assessors were blinded. Risk of bias related to missing data was high in several trials due to differential attrition, low in others because they reported data in full or did not have differential attrition, and unclear in several studies due to insufficient information. Overall, we judged the risk of selective reporting bias to be unclear because the majority of the studies were published before protocol publication became common practice.

The certainty of evidence was downgraded because of risk of bias in one outcome. For other outcomes, we downgraded the certainty of evidence due to imprecision, where there were few events, or the 95% CI was consistent with possible benefit and possible harm. We also downgraded the evidence in some places where there was heterogeneity in the direction and size of effect. Finally, we downgraded the evidence relating to costs of care because we deemed it to be imprecise, due to the lack of pooled data and effect estimates.

Potential biases in the review process

We made every attempt to reduce bias in the review process by conducting a comprehensive literature search without restrictions in terms of language or publication status. We also ensured that

we did double independent screening of search results, data extraction, risk of bias assessment and GRADE assessment, in order to reduce bias. We contacted trial authors to seek clarification and/ or obtain missing data where necessary.

In this updated review, we identified outcomes for the 'Summary of findings' tables. Two review authors were already aware of some of the trials that were included in the previously published version of the review, and this could have had an impact on the outcomes selected, resulting in a potential source of bias in this current update of the review.

Agreements and disagreements with other studies or reviews

The two other reviews that provide a quantitative synthesis on the effects of early postnatal discharge could neither confirm nor rule out the link between early discharge and neonatal and maternal morbidity and readmissions to hospital (Benahmed 2017; CETS 1997). Whilst this review found similar results for maternal readmissions within six weeks after giving birth, we found that early discharge probably slightly increases the number of infants readmitted within 28 days for neonatal morbidity. Other outcomes including maternal and infant health contacts, breastfeeding, and depression within six months align with findings from existing reviews, which conclude little to no difference between early discharge and standard length of stay. For the outcomes related to cost, we found that early discharge may slightly reduce the costs of hospital care in the period immediately following the birth up to the time of discharge (low-certainty evidence) but it may result in little to no difference in costs of postnatal care following discharge from hospital in the period up to six weeks after the birth (low-certainty evidence). This is a similar finding to the Conseil d'évaluation des technologies de la santé du Québec (CETS) review, which found that there are potential savings through early discharge only if there are no maternal or neonatal complications (CETS 1997).

Observational studies have mainly focused on the association between early discharge and infant morbidity and readmission to hospital. Our findings that early discharge probably slightly increases the number of infants readmitted within 28 days corresponds with a recent population based study by Lain and colleagues, which found that infants discharged from hospital in the first two days after birth were more likely to be readmitted for jaundice than infants who had a postnatal length of stay of three days or more (Lain 2015); although a previous study by the same author concluded that when considering all hospital related admissions (not just potentially avoidable conditions such as jaundice, which could be amenable to earlier intervention in the care pathway), a longer length of birth hospitalisation was associated with a higher risk of readmission (Lain 2014). Another recent study by Harron and colleagues found no association between postnatal length of stay and risk of readmission within 30 days for term infants. However, for premature infants between 34 weeks and 36 weeks gestation, each additional day of postnatal stay in hospital was associated with an 8.6% decreased risk of readmission (Harron 2017).

AUTHORS' CONCLUSIONS

Implications for practice

In this review, we wanted to answer the following questions in order to help women, clinicians and policymakers make informed decisions.

1. Does a policy of early postnatal discharge for healthy mothers and term infants increase infant mortality, readmissions and contacts with health services after leaving hospital?

Low-certainty evidence and wide confidence intervals mean that we are uncertain whether early discharge, compared with standard discharge has any effect on the risk of infant mortality within one year. Moderate-certainty evidence shows that early discharge, compared with standard discharge probably slightly increases infants readmitted within 28 days compared to those discharged according to standard protocol. We are uncertain whether early discharge has any effect on readmissions within seven days. Moderate-certainty evidence shows that early discharge compared with standard discharge probably makes little to no difference in the number of infants having at least one unscheduled medical consultation or contact with health professionals, although this finding must be interpreted with caution due to the considerable variation in co-interventions and support available to families at home following discharge. No trials reported on infants attending hospital casualty or emergency department within 28 days.

2. Does a policy of early postnatal discharge for healthy mothers and term infants increase maternal mortality, readmissions and contacts with health services after leaving hospital?

No studies reported on maternal mortality within six weeks postpartum, although two studies reported maternal mortality within one year, finding that no maternal deaths in either the early discharge or control group. Moderate-certainty evidence indicates that early discharge probably results in little to no difference in women readmitted within six weeks, and probably little to no difference in the number of women having at least one unscheduled contact with health professionals. The proportion of women attending hospital casualty or emergency department within six weeks after birth was not reported by any of the trials.

3. Does a policy of early postnatal discharge for healthy mothers and term infants increase breastfeeding problems, and/or decrease the duration of breastfeeding?

Moderate-certainty evidence shows that early discharge probably results in little to no difference in women breastfeeding at six weeks. When considering duration of breastfeeding, we found that there is also little to no difference between early discharge and standard discharge in the proportion of women breastfeeding at six months.

4. Does a policy of early postnatal discharge for healthy mothers and term infants influence women's satisfaction with postnatal care in hospital and following discharge?

Findings suggest that there may be little to no difference in satisfaction scores between women who were discharged early and women discharged according to standard protocol. No trials reported on women's satisfaction with their postnatal care in the community following discharge although this would likely depend



on the quality of community support available to women in the community.

5. Does a policy of early postnatal discharge for healthy mothers and term infants increase the costs of hospital and/or community postnatal care?

Low-certainty evidence suggests that early discharge may slightly reduce costs of hospital postnatal care and that early discharge may result in little to no difference in the cost of postnatal care following discharge from hospital in the six weeks after the birth. The latter is likely to depend considerably on the provision of postnatal community support available to women who are discharged early.

Implications for research

Given the limitations of the evidence to support the practice of early postnatal discharge, there continues to be a need for welldesigned trials of this intervention to inform current practice. However, future studies should be large enough to detect important differences taking into account the likelihood of attrition resulting from post-randomisation exclusions, protocol deviations (crossover) and withdrawals. The role of co-interventions in addition to early discharge is of particular importance in light of the COVID-19 pandemic, where many women and infants have not received the standard postnatal care following discharge from hospital, due to restrictions on home visiting (MacGregor 2020). Process evaluation to assess the nature and uptake of any co-interventions is of critical importance as this is likely to affect satisfaction with postnatal care and utilisation of community health services. Use of standardized approaches to outcome assessment would greatly improve the capacity to interpret results, particularly for outcomes regarding utilisation of health services following discharge from hospital, and compare the findings of future studies.

We only identified one small ongoing trial (Namagembe 2014), and in the absence of further trials (due to the difficultly of conducting them in this area of study), it is likely that observational studies would help to make the evidence more certain. Additionally, metaanalysis of individual patient data from the existing randomised trials would help to resolve some of the difficulties in analysing studies with such heterogenous interventions. Future reviews may benefit from approaching the analysis using alternative methods of comparison, based on time, to overcome the issue of heterogeneous definitions of early and late discharge.

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As part of the prepublication editorial process, this updated review has been commented on by three peers (an editor and two referees who are external to the editorial team), a member of the Pregnancy and Childbirth Group's international panel of consumers and the Group's Statistical Adviser. The authors are grateful to the following peer reviewers for their time and comments: Dr Edgardo Abalos, Vice Director, Centro Rosarino de Estudios Perinatales (CREP), Rosario, Argentina; Dr Andrew G Symon;

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CHARACTERISTICS OF STUDIES

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* Indicates the major publication for the study

MethodsDesign: RCT
Location: Maternity Hospital, Faculty of Medicine, Cairo University, Egypt
PN recruitment: "All patients were assessed after 24 h, those fulfilling our discharge criteria were ran-
domised using sealed envelopes"ParticipantsNumber of women eligible: 3786
Number of participants randomised to treatment groups: A: 1890; B 1896.
Number of post-randomisation exclusions: A: 132/1890 refused to be discharged after 24 hours; B:
188/1896 refused to be discharged after 72 hours. (all women in both groups were still eligible for early
discharge)

Number of withdrawals: A: 263/1758 lost to follow-up B 205/1708 lost to follow-up

Number of women analysed: A: 1495 B: 1503

Maternal age (years) (mean (SD)) (per group if available): A: 30.6 (6.2); B: 30.9 (5.7) Gestational age at birth (per group if available): A: 39.44 (1.47); B: 39.07 (1.28)

Inclusion criteria: caesarean section, age from 20 to 40 years; no known medical condition, e.g. diabetes, hypertension, renal, cardiac, chest problems or connective tissue disease; no obstetric complications, e.g. placenta previa, placenta accreta, accidental hemorrhage or preeclampsia; all term pregnancies 37 weeks or more; no fetal problems, e.g. intrauterine growth retardation or major congenital anomalies; and all patients who are discharged with their newborn.

Trusted evidence.	
Informed decisions.	
Better health.	

Bayoumi 2016 (Continued)	
	Exclusion criteria: age less than 20 years or more than 40 years; patients with a known medical prob- lem; patients with any obstetric complication; any preterm delivery; any fetal problem diagnosed after delivery or patients and newborns who did not meet our criteria of discharge at the 24 or 72 h interval chosen for their discharge.
Interventions	A: (n = 1890) discharge 24 hours after caesarean delivery
	B: (n = 1896) discharge 72 hours after caesarean delivery Duration of follow-up: 6 weeks
Outcomes	Wound infection
	Abdominal pain
	Secondary postpartum haemorrhage
	Successful breastfeeding
	Breast engorgement
	Intestinal ileus
	Fever above 38 C
	Stress incontinence
	Chest infection
	Deep venous thrombosis
	Pulmonary embolism
	Mood swings measure by Edinburgh Postnatal Depression Scale
	Hospital readmission
Notes	Co-interventions: "Before being discharged, all patients were given strict instructions about wound care and breast feeding. We could not offer home visits by healthcare professionals but patients could access medical help through our morning outpatient clinics or through the 24-h emergency depart- ment." Dates of study: June 2012 to February 2014.
	Funding sources: not reported
	Declarations of interest: "The authors report no declarations of interest"
Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	"randomised using sealed envelopes, containing the discharge protocol, at- tached to the file of each patient, these envelopes were opened after 24 h and patients were discharged according to the chosen protocol and those who did not fulfil the criteria of discharge after 24 h were excluded from the study" Random sequence generation not reported.
Allocation concealment (selection bias)	Low risk	"randomised using sealed envelopes, containing the discharge protocol, at- tached to the file of each patient, these envelopes were opened after 24 h and patients were discharged according to the chosen protocol and those who did not fulfil the criteria of discharge after 24 h were excluded from the study"

Bayoumi 2016 (Continued)

Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcome of hos- pital readmission
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	"All the outcome measures in the study were largely self-reported and were not confirmed by medical personnel. However, the outcome measures were not controversial and there was no reason for the participants to be dishonest in their responses"
Incomplete outcome data (attrition bias) All outcomes	Low risk	No differential attrition but no ITT analysis. 1495/1890 and 1503/1896 women were included in the analysis "All the patients came after 6 weeks for postpar- tum check-up, patients who did not attend this postpartum visit were exclud- ed from our study"
Selective reporting (re- porting bias)	Unclear risk	No indication that the trial was registered prospectively. No mention of proto- col. We do not have enough information to know if the trialists measured and reported all the outcomes that they pre-specified
Other bias	Low risk	No indication of any other sources of bias

Boulvain 2004

Study characteristics

-	
Methods	AN recruitment
	Randomisation: by telephone, using sealed envelopes. Recruitment and randomisation at > 37 weeks' gestation.
	Blinding: caregivers, women and outcome assessment unblinded.
	Follow-up to 6 months pp.
	Analysis: intention to treat.
	Loss to follow-up: 16/459 = 3.5% comprising 11 in early discharge arm and 5 in comparison arm.
	Duration: November 1998 to October 2000.
	Setting: urban tertiary level hospital, Switzerland.
Participants	Number of women eligible: 2324 Number of participants randomised to treatment groups: 459 (A: 228; B: 231) Number of post-randomisation exclusions: A: 1; B: 0. Number of women who remained in trial: A: 227; B: 231. Number of women analysis: A: 227; B: 231.
	Inclusion criteria: primiparous and multiparous women at low risk of caesarean delivery and/or post- natal complications > 37 weeks' gestation.
	Exclusion criteria: women with a strong preference for long or short length of stay; placenta praevia; pre-eclampsia; diabetes treated with insulin; medical complications of pregnancy requiring postna- tal surveillance; past history of postnatal complications (e.g. postnatal depression); difficult socio-eco- nomic situation; multiple pregnancy; suspected intrauterine growth retardation or large infant for ges- tational age; fetal malformation or genetic disease.
	Characteristics: maternal age: A: mean 29 years (SD 4.8); B: mean 29 years (SD 5.5); primiparous A: 60%; B: 57%; married A: 83%; B: 82%; income < 50,000 CHF A: 27%; B: 24%; tertiary education A: 48%; B: 49%;



3oulvain 2004 (Continued)	Swiss origin A: 31%; B: (SD 405).	30%; current smoker A: 25%; B: 17%, infant birthweight A: 3420 (SD 435); B: 3480	
Interventions	A: (n = 228): home based postnatal care with discharge planned for 24 < 48 hours following vaginal births and 72 < 84 hours after caesarean.		
	· · · · ·	ased postnatal care with discharge planned 4 to 5 days postpartum (pp) follow- 6 to 7 days pp following caesareans.	
	Co-intervention (A and B): minimum of 2 nurse home visits and 10 phone calls; number and timing de- termined by the family.		
Outcomes	Infant readmissions wi	thin 28 days and in first 6 months pp.	
	Maternal readmissions	within 28 days and in first 6 months pp.	
	Proportion of women c	lepressed at 28 days pp.	
	Proportion of women b	preastfeeding at 7 days and 28 days pp.	
	Proportion of women reporting breastfeeding problems.		
	Maternal satisfaction with postnatal care.		
	Costs of hospital care post birth to discharge.		
	Costs of postnatal care post discharge to 6 weeks pp.		
Notes	Significant non-compliance in A group; mean length of stay A: 65 hours; B: 106 hours.		
	Dates of study: November 1998 to October 2000		
	Funding sources: "The study received financial support from the Swiss National Science Foundation (grant #32-52954.97) and from the Quality of Care Program, Geneva University Hospitals."		
	Declarations of interest: not reported		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	"The allocation sequence was prepared with a computer program generating a random sequence of numbers in blocks of varying size (four, six and eight participants), arranged in random order"	
Allocation concealment (selection bias)	Low risk	"Consenting women were allocated to home or hospital care postpartum by opening a consecutively numbered, opaque, sealed envelope. The envelopes were opened by a third party, usually the trial co-ordinator during a telephone call with the research midwife"	

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"The allocation sequence was prepared with a computer program generating a random sequence of numbers in blocks of varying size (four, six and eight participants), arranged in random order"
Allocation concealment (selection bias)	Low risk	"Consenting women were allocated to home or hospital care postpartum by opening a consecutively numbered, opaque, sealed envelope. The envelopes were opened by a third party, usually the trial co-ordinator during a telephone call with the research midwife"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcome of hospital readmission
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Unclear for outcome assessors.
Incomplete outcome data (attrition bias)	Low risk	Low attrition and not differential



Boulvain 2004 (Continued) All outcomes

Selective reporting (re- porting bias)	Unclear risk	Not common practice to publish protocol at that time.	
Other bias	Low risk	Nothing to indicate any other source of bias	

Brooten 1994

Study characteristics	
Methods	Method of randomisation: sealed envelopes, recruitment and randomisation 24 hours post caesarean.
	Blinding: caregivers and women unblinded, blinding of outcome assessment unclear.
	Loss to follow-up: not reported
	Analysis: by intention to treat.
	Follow-up: 8 weeks pp.
	Duration: August 1988 to January 1991.
	Setting: urban tertiary-level hospital, US.
Participants	Number of women eligible: not reported
	Number of participants randomised to treatment groups: not reported
	Number of post-randomisation exclusions: not reported.
	Number of withdrawals: not reported
	Number of women analysed (per group if available): A: 61; B: 61.
	Inclusion criteria: unplanned caesarean delivery in hospital, English speaking; healthy mother and in- fant (range: 2270 to 4680 g; 36 to 43 weeks' gestation).
	Characteristics:
	Maternal age: early mean 29 (SD 6), standard 28 (SD 6); Marital status: early 67% married, standard 56% married; Income (< \$10,000) early 29%, standard 33%. Education (< high school) early 15%, standard 21%. 'Race' (African American and 'non-white') early 53%, standard 61%. Birthweight: early mean 3305 g (SD 483), standard 3440 g (SD 572). Gestation: early mean 39 weeks (SD 1.5), standard 39 weeks (SD 1.7).
Interventions	A: discharge 'earlier than usual' (mean stay of 3.6 days);
	B: discharge according to 'routine hospital practice' (mean stay of 4.8 days). Co-interventions: A had minimum of 2 home visits post discharge, plus 10 phone calls to 8 weeks, plus women had phone number to nurse and physician. B had no routine follow-up care at home post discharge.
Outcomes	Infant and maternal re-admissions. Infant and maternal acute care visits. Maternal satisfaction with care. Proportion of infants immunized. Mean cost per woman for care from birth to discharge and for post discharge care.



Brooten 1994 (Continued)NotesWomen in standard care considered ready for discharge if: ambulatory, voiding, tolerating normal di-
et, passing flatus, experiencing normal uterine involution, afebrile for 24 hours, uncomplicated wound
healing, removal of sutures and an adequate blood count. Women in early group had same criteria ex-
cept: staple removal and afebrile status for at least 24 hours.If women in early group did not meet these criteria, they were not discharged early but still received co-
interventions and were analysed with early group.Dates of study: August 1988 to January 1991Funding sources: "This study was supported by a grant (Pol-NR-01859) from the National Institute for
Nursing Research, National Institutes of Health"

Declarations of interest: not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Randomisation list prepared in advance by a statistician.
Allocation concealment (selection bias)	Low risk	Adequate.Sequence of sealed envelopes containing cards with the name of the next treatment group
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcome of hos- pital readmissions
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Unclear for outcome assessors.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up not reported.
Selective reporting (re- porting bias)	Unclear risk	Not common practice to publish protocol at that time
Other bias	Low risk	Nothing to indicate any other source of bias

Burnell 1982

Methods	Design: 4-arm study, 2 arms allocated by randomisation, 2 arms allocated according to woman's prefer ence AN recruitment
	Setting: Oxford, UK
Participants	Number of women eligible: "242 women were considered at booking for inclusion in the study" Number of participants randomised to treatment groups: A: 23; B: 19. (only those who did not express a preference for short or long stay in hospital were allocated randomly) Number of post-randomisation exclusions: not reported

Burnell 1982 (Continued)		
	in early discharge grou Number of withdrawals Number of women ana Maternal age (years) (m Gestational age at deliv	p remained in trial: 153 (106 participants were self-selecting not randomised: 33 p, 73 in standard stay group) s: not reported lysed (per group if available): 148 nean (SD)): not reported <i>y</i> ery (weeks): A: 39.44 (1.47); B: 39.07 (1.28) Inclusion criteria: All women over 18 ckground and social class were included.
	Exclusion criteria: wom	en with major clinical disorders and those to be delivered by caesarean section
Interventions	A: (n = 23) transfer to pe	ostnatal care at home 48 hours after delivery
	B: (n = 19) conventiona Duration of follow-up: (l hospital stay of 8 to 9 days 6 weeks
Outcomes	Physical problems of m	nother and child
	Feeding and related pr	oblems
	Emotional state of the	mother
	Mother's attitude towa	rds her ability to cope, to her baby and to postnatal care
Notes	Co-interventions: not reported Dates of study: not reported Funding sources: not reported Declarations of interest: not reported	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	"Allocated to either (c) early discharge: randomised. or (d) usual stay: ran-
· · · · ·		domised."
		domised." No details reported about how randomisation was done
Allocation concealment (selection bias)	Unclear risk	
Allocation concealment	Unclear risk High risk	No details reported about how randomisation was done
Allocation concealment (selection bias) Blinding of participants and personnel (perfor- mance bias)		No details reported about how randomisation was done Not reported Not possible to blind women. Lack of blinding could have an effect on out-
Allocation concealment (selection bias) Blinding of participants and personnel (perfor- mance bias) All outcomes Blinding of outcome as- sessment (detection bias)	High risk	No details reported about how randomisation was done Not reported Not possible to blind women. Lack of blinding could have an effect on out- comes.
Allocation concealment (selection bias) Blinding of participants and personnel (perfor- mance bias) All outcomes Blinding of outcome as- sessment (detection bias) All outcomes Incomplete outcome data (attrition bias)	High risk Unclear risk	No details reported about how randomisation was done Not reported Not possible to blind women. Lack of blinding could have an effect on out- comes. Not reported Not reported No data reported per randomised group so we do not know how many women
Allocation concealment (selection bias)Blinding of participants and personnel (perfor- mance bias) All outcomesBlinding of outcome as- sessment (detection bias) All outcomesIncomplete outcome data (attrition bias) All outcomesIncomplete reporting (re-	High risk Unclear risk Unclear risk	No details reported about how randomisation was done Not reported Not possible to blind women. Lack of blinding could have an effect on outcomes. Not reported Not reported Not reported Not reported



Study characteristics			
Methods	Method of randomisation: sealed envelopes placed on file prior to home visit at 38 weeks. Recruitment and randomisation at 37 weeks' gestation.		
	Blinding: caregivers, women and outcome assessment unblinded.		
	Follow-up to 1 month pp (most outcomes).		
	Loss to follow-up: 58/189 (30.7%), including 46 post-randomisation exclusions, 10 not compliant with randomisation outcome, 2 withdrawals; 97/131 returned 1 month questionnaires.		
	Analysis: not intention to treat.		
	Duration: not reported.		
	Setting: urban tertiary-level hospital, Canada.		
Participants	Number of women eligible: 189		
	Number of participants randomised: 189 (not reported per group)		
	Number of post-randomisation exclusions: 45 no longer met the inclusion criteria		
	Number of withdrawals: 1 home birth, 2 reversed their decision to participate, 10 chose to go home at times other than the discharge times to which they had been assessed.		
	Number of women analysed: A: 44; B: 49. C 38		
	Inclusion criteria: normal labour and hospital birth. Exclusion criteria: caesarean, forceps delivery. Participants were 53% primiparous; mean maternal age 30.2 (SD 3.8); 93% married or living with part- ner; 58% combined family income > \$40,000; 65% completed junior college or university; 95% 'Cau- casian'; mean paternal age 32.9 (SD 5.5). No significant group differences found on demographic characteristics (but no data provided by group).		
Interventions	3 groups, discharge at: 12 to 24 hours (n = 44), 25 to 48 hours (n = 49) or 4 days (n = 38).		
	Co-interventions:		
	 12 to 24 hrs - 1 home visit by nurse antenatally; 5 home visits post discharge; 25 to 48 hours - 1 antenatal home visit; 3 home visits post discharge; 4 days - 1 home visit antenatally; 1 home visit post discharge. 		
Outcomes	Infant and maternal re-admissions. Maternal depression, anxiety and confidence. Maternal satisfaction with nursing care. Breastfeeding at 1 month pp. Referrals to physicians for maternal and infant health issues.		
Notes	Mean length of stay:		
	12 to 24 hrs:1.12 days (SD 0.4). 25 to 48 hrs: 2.06 days (SD 0.6). 4 days: 4.03 days (SD 0.7).		
	Study nurses participated in 2 weeks special training for the early discharge program.		
	Dates of study: not reported		
	Funding sources: not reported		



Carty 1990 (Continued)

Declarations of interest: not reported

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Using a table of random numbers.
Allocation concealment (selection bias)	Low risk	Sealed opaque envelopes containing the assignment was placed on the file of each prospective participant and opened by the nurse.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcome of hospital readmissions
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Unclear for outcome assessors.
Incomplete outcome data (attrition bias) All outcomes	High risk	Differential attrition (82%, 74% and 66% returned follow-up questionnaires)
Selective reporting (re- porting bias)	Unclear risk	Not common practice to publish protocol at that time
Other bias	Low risk	Nothing to indicate any other source of bias

Chiong 2012

Study characteristic	s
Methods	Design: RCT PN recruitment: "The envelope was opened on post cesarean day 1 morning ward round by the health care provider to reveal the allocation" Location: A university hospital in Kuala Lumpur, Malaysia
Participants	Number of women eligible: 360 Number of participants randomised to treatment groups: 360 (A: 179; B: 181) Number of post-randomisation exclusions: A: 9/179; B: 9/181 Number of women who remained in trial: A: 170; B: 172 Number of withdrawals: A: 28/170 (9 declined discharge, 19 either mother or baby not fit for discharge) B: 24/172 (12 took discharge on day 1, 12 either mother or baby not fit for discharge) Number of women analysed: A: 142; B: 148.
	Maternal age (years) (mean (SD)): A: 31.8 (4.6); B: 31.5 (4.1)
	Gestational age at birth (per group if available): A: 38.8 (1.0); B: 38.7 (0.9)
	Inclusion criteria: planned caesarean delivery, age 18 years or older, and gestation 37 weeks or greater with a singleton pregnancy
	Exclusion criteria: grossly abnormal fetus, established medical problems, or operative factors likely to preclude day 1 discharge (e.g., pre gestational diabetes, preeclampsia, epilepsy, cardiac disease, renal disease, connective tissue disease, antiphospholipid syndrome, 2 or more previous caesarean deliver- ies, and major previa).

Chiong 2012 (Continued)	
	Protocol predefined discharge criteria which were; surgically uncomplicated caesarean, blood loss 800 mL or less, post operative haemoglobin of 80 g/L or greater, unassisted ambulation to the bathroom, established urination, tolerated at least 1 normal meal, afebrile (temperature less that 38 ^o C in the last 24 hours), normal blood pressure and healthcare professional willing to discharge. Neonatal discharge was predicated on standard paediatric practice.
	Characteristics: similar between the arms for maternal age (31.8 versus 31.5 years); gestational age 38.8 versus 38.7 weeks; parity both 1; ethnicity; Malay 104 (58.1%) versus 100 (55.2%), Indian 29 (16.2%) versus 43 (23.8%), Chinese 35 (19.6%) versus 28 (15.5%), other 11 (6.1%) versus 10 (5.5%). Employed: 134 (74.9%) versus 137 (75.7%), Homemaker or student 45 (25.1%) versus 44 (24.3%); Education: primary 9 (5.0%) versus 2 (1.1%), secondary 67 (37.4%) versus 76 (42%), and Tertiary 103 (57.5%) versus 103 (56.9%).
Interventions	A: (n = 179) discharge 1 day after caesarean B: (n = 181) discharge 2 days after caesarean Duration of follow-up: 6 weeks
Outcomes	Primary outcomes: satisfaction with discharge protocol at 2 weeks after discharge by means of a 5 point Likert scale (top 2 responses taken as demonstrating satisfaction and other 3 responses as non satisfaction) and self reported breastfeeding at 6 weeks.
	Secondary outcomes assessed at 2 weeks:
	Double check to see and add readmissions
	 satisfaction with discharge protocol based on the full 5-point Likert scale, general well-being score using a 10-point numerical rating scale, recommendation of their timing of discharge after caesarean delivery to a friend (5-point Likert scale), preferred length of hospital stay after caesarean delivery, newborn feeding status, maternal or newborn unscheduled medical consultation, maternal antibiotics and caesarean wound condition (observed or self-report of skin reddening around the wound, purulent wound discharge, or wound gaping of any severity). Secondary outcomes assessed at 6 weeks after discharge: maternal and infant rehospitalization general well-being score, maternal or newborn unscheduled medical consultation, maternal or newborn unscheduled medical consultation, general well-being score, maternal or newborn unscheduled medical consultation, general well-being score, maternal or newborn unscheduled medical consultation, maternal or newborn unscheduled medical consultation, maternal antibiotic, caesarean wound condition, and assessment of maternal anxiety and depression using the Hospital Anxiety and Depression scale. Participants were subsequently also contacted to establish whether readmissions within the first 6 weeks after discharge had occurred for mother or newborn.
Notes	Co-interventions: "Home visit by health care professionals from our center were not on offer but our patients have free access to our clinics and acute care facilities, which were opened at all hours. An appointment was routinely given for maternal and newborn follow-up at 2 weeks after caesarean delivery. After this, another routine appointment would be given at 6 weeks postdelivery. Additional appointments may be scheduled as clinically required"
	Dates of study: November 2010 to February 2012
	Funding sources: University of Malaya
	Declarations of interest: "Financial Disclosure: The authors did not report any potential conflicts of in- terest."

Risk of bias



Chiong 2012 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"online random number generator (www.random.org) in random blocks of 4 to 8 on a 1:1 ratio"
Allocation concealment (selection bias)	Low risk	"A sealed numbered opaque envelope containing the allocated discharge protocol was then attached to the chart of each participant. The envelope was opened on post cesarean day 1 morning ward round by the health care provider to reveal the allocation"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcome of hos- pital readmissions
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	No differential attrition; the analysis included data from women who did not adhere to the allocated group and from women who were no longer eligible for early discharge but they did not ana lyze data from women who no longer met the trial eligibility criteria. Overall level of missing data for any reason: 5%
Selective reporting (re- porting bias)	Unclear risk	No protocol available, retrospectively registered. Insufficient information to judge whether all pre-specified outcomes were reported
Other bias	Low risk	Nothing to indicate any other source of bias

Gagnon 1997

Study characteristic	s
Methods	Randomised controlled trial
	Antenatal recruitment
	Setting: urban university hospital, Canada.
Participants	Number of women eligible: 938 Number of participants randomised to treatment groups: 360 (A: 183; B: 177) Number of post-randomisation exclusions: A: 85/183; B: 74/177. Number of women who remained in trial: A: 80; B: 100. Number of withdrawals: A: 18/98; B: 3/103 (no explanations given for these withdrawals) Number of women analysed: A: 78; B: 97. Inclusion criteria at randomisation: parity 0 to 4; normal pregnancy (no medical conditions, not breech); English, French or Spanish speaking. intrapartum exclusion criteria were rupture of membranes> 24 hours before birth, caesarean deliv- ery, intrapartum blood loss > 500 mL, and third- or 4th-degree perineal laceration. Postpartum exclu- sion criteria for the mother were soft fundus with excessive bleeding, inability to void adequately, in- ability to ambulate easily or to care for herself and her infant, non receipt of RhoGAM when indicated, and medical conditions requiring close supervision; for the infant they were multiple birth, birth weight < 2500 g, gestational age < 36 completed weeks, abnormal newborn physical examination, and intoler- ance to feedings (breast or bottle) in hospital. Characteristics of participants:



Gagnon 1997 (Continued)	
	Mean maternal age (SD): early 29.6 (4.7), standard 29.1 (5.3). Parity (% nulliparous): early 38%, standard 34%. Living with a partner: early 85.5%, standard 93.8%. % 'blue collar': early 21.8%, standard 16.5%. Mean years of maternal education (SD): early 13.8 (3.8), standard 14.0 (3.9). % recent immigrants: early 14.7%, standard 24.7%. Mean birthweight (SD): early 3389g (419), standard 3496g (364). Mean gestation (SD): early 39.3 (1.3), standard 39.5 (1.1). Planned to breastfeed: early 70.5%, standard 54.6%. Smoking in pregnancy: early 23.1%, standard 9.3%.
Interventions	A: (n = 183 randomised, 98 remained eligible after giving birth) "The early postpartum discharge pro- gram consisted of a postpartum hospital stay of 6 to 36 hours and nursing care by telephone, within 48 hours pp and at 10 days pp, and also at home, at 34 to 38 weeks' gestation and at 3 and 5 days pp. 2 part-time nurses made the visits, which lasted a maximum of 1 hour. The postnatal intervention was of- fered only to women who had left the hospital within 36 hours of the birth to encourage them to leave the hospital early if they had been randomised into this group and to maximize the generalisability of the intervention" B: (n = 177 randomised, 103 remained eligible after giving birth) postpartum stay of 48 to 72 hours and follow-up as determined by the woman's and infant's physicians. Duration of follow-up: 1 month
Outcomes	Breastfeeding Infant health contacts post discharge. Maternal satisfaction with care to day 10 pp. Perceived maternal competency.
Notes	Significant non-compliance with early discharge allocation in the early group - mean length of stay 37.5 hours (26 women - 33% went home later than planned).
	Dates of study: January to December 1990
	Funding sources: "Funded by a grant from Le Fonds de la Recherche en Santé du Québec. A.J.G. was supported by the Medical Research Counil and the National Health Research and Development Pro- gram of Canada"
	Declarations of interest: not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Subjects were stratified by date of immigration and parity and randomised 32-38 weeks' gestation in blocks of 8 according to a random number table
Allocation concealment (selection bias)	Low risk	Group assignment was placed in sealed opaque envelopes.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	No blinding and the outcome is likely to be influenced by lack of blinding
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Unclear for outcome assessors.
Incomplete outcome data (attrition bias)	High risk	Differential withdrawals across groups: A: 18/98; B: 3/103 (no explanations given for these withdrawals)



Gagnon 1997 (Continued) All outcomes

Selective reporting (re- porting bias)	Unclear risk	Not common practice to publish protocol at that time	
Other bias	Low risk	Nothing to indicate any other source of bias	

Hellman 1962

Study characteristics	
Methods	Postnatal recruitment
	Blinding: caregivers, women and outcome assessment unblinded.
	Follow-up to 3 weeks pp.
	Loss to follow-up: not reported.
	Analysis: by intention to treat.
	Duration: 1 July 1959 to 30 June 1960.
	Setting: urban hospital, New York, US.
Participants	Number of women eligible: not reported
	Number of participants randomised to treatment groups: 2257 (A: 1941; B: 316) Number of post-randomisation exclusions: not reported
	Number of women analysed: A: 1941; B: 316 Inclusion criteria: hospital birth, mothers deemed eligible for early discharge, babies predominantly > 2500g, baby gestation not specified.
	Exclusion criteria: caesarean section, stillbirth, no English. Characteristics of participants:
	Median age: early 23.6 yrs, standard 23.8 yrs. No living children: early 28%, standard 29%. Married: early 70%, standard 73%. Welfare/no income: early 16%, standard 13%. Ethnicity: Negro/Puerto Rican early 81%, standard 85%. Few details available on babies.
Interventions	A: (n = 1941): discharge before 72 hrs. B: (n = 316): discharge after 5 days.
	Co-interventions: midwife home visits post discharge (3 in A, 2 in B) for examination of mother and ba- by and other data collection (not for 'helping' mothers).
	Duration of follow-up: 3-4 weeks after discharge
Outcomes	Infant and maternal re-admissions within 3 weeks. Infants deaths. Proportion of women breastfeeding at 3 weeks. Reported infant feeding problems. Proportion of women dissatisfied with hospital postnatal care. Proportion of women dissatisfied with length of stay. Proportion of fathers thinking stay too short.



Hellman 1962 (Continued)

	Proportion of fathers thinking stay too long. Limited cost data.	
Notes	Dates of study: not reported	
	Funding sources: 'United States Public Service Grant'	
	Declarations of interest: not reported	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	High risk	"Included in the experimental group was every patient who was discharged from the hospital within 72 hours during a five-day work week" control group: "selected by numbering every patient delivered in the hospital and choosing the patients who were to be in the control group by a series of previously ran- domised numbers"
Allocation concealment (selection bias)	Unclear risk	Insufficient information to permit judgement of 'low risk' or 'high risk'
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcome of hospital readmissions
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Unclear for outcome assessors.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up not reported.
Selective reporting (re- porting bias)	Unclear risk	Not common practice to publish protocol at that time
Other bias	Low risk	Nothing to indicate any other source of bias

Kruse 2021

Study characteristic	S
Methods	2-arm, parallel assignment RCT
	No blinding
	Antenatal recruitment
	Setting: Department of Obstetrics and Gynecology, Herning Hospital, Denmark
Participants	Number of women eligible: 328
	Number of participants randomised to treatment groups: 143 (A)72 (B)71)
	Number of post-randomisation exclusions (per group if available): A) 0 B) 0

Number of women included in ITT analysis: A) 72, B) 71 Number of women responding to questionnaire (Parents' Postnatal Sense of Security): A) 53/72 B)
 52/71 Maternal age (years) (mean (SD)) (per group if available): A) 33.4 (4.5) B) 32.6 (4.6) Gestational age at birth (per group if available):38.7 (SD 0.54) 38.9 (SD 0.48) Inclusion criteria: planned caesarean section Exclusion criteria: nulliparity, multiple pregnancy, gestational age < 37 + 0, planned prolonged postoperative observation, pre-pregnancy BMI > 35 kg/m2, maternal age,18 yrs, living alone, or inability to read and write Danish. Discharge criteria: no symptoms of postpartum complications, sufficient mobilisation and analgesia, normal voiding function and successfully initiated breastfeeding if anticipated.
Inclusion Criteria:
 18 years or older Planned elective caesarean of multiparous women Singleton pregnancy Gestational age at least 37 + 0 weeks Pre pregnancy BMI < 35
Exclusion Criteria:
 Lack of consent Women with no or little understanding of and ability to speak Danish Expected maternal or neonatal complications after delivery
A: (n = 72) [discharge at 28 hours] B: (n = 71) [discharge > 48 hours] Duration of follow-up: 1 month
Primary outcome:
• Parents' Postnatal Sense of Security at 1 week after delivery, using validated PPSS-questionnaire
Secondary outcomes:
 Pain during the first week postpartum (using numeric rating scale) Use of analgesia during the first week postpartum (Daily consumption in mg of paracetamol, NSAII and opioid.)
 Mobilization at 5 days postpartum (Step count measured by an activity monitor (Fitbit Flex)) Breastfeeding at 6 months after delivery (Duration and extent of breastfeeding)
 Readmissions at 28 days postpartum (Number of readmissions and length of hospital stay in numbe of days)
 Surgical complications at 28 days postpartum (Number of complications using the Clavien-Dindo Classification (1-5))
 Complications in the postnatal period (28 days postpartum) (Number and type of complications using the ICD-10 classification)
 Contacts to the health care system at 28 days postpartum (Number of contacts and site of contact (primary or secondary care))
Co-interventions: home visit by a midwife 48-72 hours after delivery Dates of study: September 2016 - September 2019 Funding sources: Brodene Hartmanns Foundation Declarations of interest: the authors have 'no conflicts of interest to declare'
Data and additional information provided by author (June 2020)



Kruse 2021 (Continued)

 "Randomization was conducted using a computerized random number generator (www.randomization.com). The randomisation was generated in a 1:1 ratio in blocks of two and four and stratified according to site and BMI < 30 or 30 kg/m2" "Envelopes were sealed individually and opened by the woman when written consent was obtained."Correspondence from study author: "sequentially numbered, opaque sealed envelopes" Blinding not possible for mothers but unlikely to influence the outcome of hospital readmissions
ten consent was obtained."Correspondence from study author: "sequentially numbered, opaque sealed envelopes" Blinding not possible for mothers but unlikely to influence the outcome of hos-
Correspondence from study author: "the outcome assessors were not blinded to the allocations"
Very low attrition: 7/72 and 5/71 excluded from per-protocol analysis due to emergency caesarean, vaginal delivery or decision not to adhere to allocated group but these women were included in ITT analysis). All women included in the ITT analysis.
No published protocol. Trial was registered prospectively and not all pre-spec- ified outcomes are reported in full but correspondence with trial author con- firms that the outcomes were measured and the data analysis is in process.
No indication of other sources of bias

McKeever 2002

Study characteristic	S
Methods	Design: RCT Setting: Toronto, Canada PN recruitment
Participants	Number of women eligible: 214 Number of participants randomised to treatment groups: 148 (A: 72; B: 66) Number of post-randomisation exclusions (per group if available): A: 1/72; B: 0/66. Number of withdrawals: A: 7 withdrew/lost to follow-up, 5 excluded from analysis; B: 11 withdrew/lost to follow-up, 7 excluded from analysis Number of women analysed: A: 56; B: 36.
	Maternal age (years) (mean, SD): A: 32.0 (4.2); B: 33.1 (4.4)
	Gestational age at delivery (weeks): A: 38–40 wk 92%, 41 wk 8%; B: 38–40 wk 81%, 41 wk 19%
	Inclusion criteria: Mothers were eligible if they had delivered a live, singleton infant within the pre- ceding 12 hours, were at least 21 years of age, resided in the defined metropolitan area, had a tele- phone, intended to breastfeed, were breastfeeding at discharge, and would receive satisfactory sup- port at home. Satisfactory support was determined by postpartum nurses who assessed mothers' cir- cumstances to ensure the safety and well-being of infants and mothers. The main criterion was that mothers were not isolated, and had accessible family and friends to provide assistance when neces- sary. Their newborns were eligible to participate if they were 35 weeks' gestational age or greater, wer

McKeever 2002 (Continued)	breastfed at discharge, and did not have congenital anomalies or morbidities, including hyperbilirubi- naemia, blood group incompatibility, or sepsis. Exclusion criteria: Women were excluded if they did not speak English and had experienced caesarean
	deliveries, postpartum complications, and morbidities such as fever and abnormal bleeding, chronic illnesses, or disabilities.
Interventions	A: (n = 53) discharged at 24 to 36 hours postpartum and scheduled to receive up to 3 home visits from community nurses qualified as lactation consultants. B: (n = 48) "standard length of hospitalisation" Duration of follow-up: 12 days postpartum
Outcomes	Proportion of baby's feeds in past 24 hr that were exclusively breastfeeding
	% of mothers who used no supplementation in past 24 hr
	Length of hospitalisation
	Adverse infant outcomes between the term standard care and experimental groups, such as hyper- bilirubinaemia, dehydration, or weight loss
Notes	Co-interventions: "As a matter of standard hospital care, mothers of near-term infants in both groups were made aware of the outpatient hospital breastfeeding clinic, and were encouraged to use a pre ex- isting 24-hour telephone help line. Mothers of term infants in both groups were also provided with in- formation on these services." Dates of study: July 1999 to December 2000
	Funding sources: "This project was supported by a grant from the Health Transition Fund, Canada,Ot- tawa,and by a financial contribution from The Hospital for Sick Children Foundation,Toronto,Ontario, Canada." Declarations of interest: not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"Using central randomisation procedures, eligible mother-newborn pairs were stratified as term or near term (35–37 weeks' gestational age), and allocated to either the standard or experimental groups by research staff who had no con- tact with patient"
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Not possible to blind women. Lack of blinding not like to have an effect on the outcomes measured in the trial.
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	"Interviewers were originally blinded to group status. However, in the course of answering questions about postpartum care and satisfaction, mothers inad- vertently revealed their group status."
Incomplete outcome data (attrition bias) All outcomes	High risk	Slightly higher attrition in early discharge group. In the full paper, analysis is carried out only in women who completed follow-up but abstract seems to report outcomes for the full number of women randomised.
Selective reporting (re- porting bias)	High risk	Outcomes not reported in full per randomised group. No protocol (study pub- lished before common practice of publishing protocol)



McKeever 2002 (Continued)

Other bias

Low risk

Nothing to indicate any other source of bias

Sainz Bueno 2005

Study characteristics	
Methods	Recruitment and randomisation in postnatal ward.
	Blinding: caregivers and women unblinded to intervention; blinding outcome assessment unclear.
	Follow-up to 6 months pp.
	Analysis by intention to treat.
	Loss to follow-up: 36/430 (8.4%), including 22 women who did not attend 7 to10 days pp follow-up; 14 women withdrew consent, see note regarding missing data for maternal satisfaction.
	Duration: April 1999 to April 2001.
	Setting: urban maternity hospital, Spain.
Participants	430 women recruited and randomised; 213 to early discharge and 217 to standard care
	Number of withdrawals: A: 19; B: 18. Number of women analysed (per group if available): A: 213; B: 217
	Inclusion criteria: primiparous and multiparous women deemed eligible for early discharge, => 37 weeks' gestation with baby of appropriate weight for gestational age; vaginal birth; residence within 20 km of the hospital.
	Characteristics: age > 30 years A: 41.8%; B: 41.1%; primiparous A: 36.6%; B: 37.8%; married A: 97.2%; B: 97.2%; completed secondary education A: 22.5%; B: 14.7%; infant birthweight A: 3348 g (SD 396); B: 3335 g (SD 372); gestation A: 39.5 weeks (SD 1.13); B: 39.5 weeks (SD 1.12); spontaneous vaginal birth A: 87.8%; B: 88.5%.
Interventions	A: (n = 213): discharge planned for < 24 hours.
	B: (n = 217): minimum stay of 48 hours.
	Co-intervention: A monitored at home for first 24 to 48 hours post discharge by qualified nurse; A and B attended visit in clinic at 7 to 10 days pp.
Outcomes	Infant readmissions in first 28 days.
	Maternal readmissions in first 28 days.
	Proportion depressed at 4 weeks pp.
	Proportion breastfeeding at 4 weeks, 12 weeks and 6 months pp.
	Proportion reporting maternal exhaustion at 7 to 10 days and 4 weeks pp.
	Proportion reporting physical health problems in first 6 months.
	Maternal satisfaction with postnatal care.
	Proportion of women saying length of stay too long.
	Costs of hospital care post birth; costs of maternal and infant readmissions; cost of maternal and neonatal consultations (post discharge).



Sainz Bueno 2005 (Continued)

Librarv

Significant missing data for maternal satisfaction, with differential loss to follow-up I 17.8% missing, C 42% missing data.

Dates of study: April 1999 to April 2001

Funding sources: not reported

Declarations of interest: not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	"randomisation by blocks"
		Insufficient information in the report to assess whether sequence generation was adequate.
Allocation concealment (selection bias)	Low risk	"opaque sealed envelopes"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcome of hos- pital readmissions
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Unclear for outcome assessors.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No details about how missing data was accounted for
Selective reporting (re- porting bias)	Unclear risk	Not common practice to publish protocol at that time
Other bias	Low risk	Nothing to indicate any other source of bias

Smith-Hanrahan 1995

Study characteristi	cs
Methods	Method of randomisation: unclear. Recruitment and randomisation: on admission to postnatal unit post birth.
	Blinding: caregivers and women unblinded; outcome assessment - blinding unclear.
	Follow-up to 6 weeks pp.
	Loss to follow-up: 44/125 (35.2%); all post-randomisation exclusions.
	Analysis: not intention to treat.
	Duration: not reported.
	Setting: tertiary teaching hospital, eastern Canada.



Risk of bias		
	Meidcine of McGill University" Declarations of interest: Not reported	
	Funding sources: "This research was supported by a grant from the Fraser Fund of the Royal Vistoria Hospital and a computer equipment grant form the Faculties of Graduate Studies and Research and	
	Dates of study: Not reported	
Notes	29 women allocated to standard care were sent home early due to bed shortages. The authors ana lyze outcomes for this group separately from the early and standard care groups. For the purposes of this review analyses have been done re-combining these 29 women with the standard care group, to ap- proximate more closely an intention-to-treat analysis.	
Outcomes	Infant and maternal re-admissions to 6 weeks pp. Proportion of women breastfeeding at 6 weeks. Maternal fatigue intensity score at 2 to 3 days pp, 1 week pp and 6 weeks pp.	
	Co-inteventions: No antenatal preparation for discharge in either group. Early group received telephone call from nurse within 24 hours of discharge leading to a decision to vis it or continue to consult by phone; also received phone number for postnatal follow-up service which could be called at any time; followed by usual paediatric and obstetric office visits. Standard discharge group received traditional follow-up post discharge - visit to paediatric office at 2 weeks and obstetric office visit at 6 weeks.	
Interventions	A: (n = 58): early discharge: before 60 hrs. C (n = 57): discharge: after stay of 60 hrs or more.	
	Maternal age: early mean 29.5 (SD 4.5), standard mean 29.3 (SD 4.63). Parity: early 37.1% primiparous, standard 58.7% primiparous. Marital status: early 97.1% married, standard 95.7% married. All vaginal births. Income - % > 40,000+: early 74.1%, standard 55%. Completed college/university education: early 73.5%, standard 54.6%. % not of Canadian/US nationality: early 39.5%, standard 23.5%.	
	Inclusion criteria: English or French speaking; another adult present at home at least 12 hours/day for 1st 2 days post discharge; no major obstetrical complications at any stage; no prolonged mother-infan separation in hospital (24 hrs+); medical follow-up plan before discharge; no complications in infant: 2500-4500 g; good colour/activity level; vital signs normal; voided and stooled; feeding established. Characteristics of participants:	
	B: 21/67 (6 did not complete the study, 7 did not stay long enough, 6 received postnatal home fol- low-up, 2 had to stay longer in hospital, 29 transferred to early discharge group due to bed shortages) Number of women analysed (per group if available): A: 35 B: 17	
	A: 23/58 (3 did not complete the study, 20 excluded - 18 due to jaundice, 1 due to Down's syndrome, 1 due to mother's request)	
	Total number of withdrawals:	
	Number of participants randomised to treatment groups: A: 58; B: 67 Number of post-randomisation exclusions: A: 20; B: 29	
Participants	139 women approached; 125 agreed and randomised	

Smith-Hanrahan 1995 (Continued)

Random sequence genera- tion (selection bias)	Unclear risk	Insufficient information in the report to assess whether sequence generation was adequate.
Allocation concealment (selection bias)	Unclear risk	Not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcome of hos- pital readmissions
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Unclear for outcome assessors.
Incomplete outcome data (attrition bias) All outcomes	Low risk	High number of withdrawals (due to participants no longer meeting trial cri- teria) but not differential. A: 23/58 (3 did not complete the study, 20 excluded - 18 due to jaundice, 1 due to Down's syndrome, 1 due to mother's request); B: 21/67 (6 did not complete the study, 7 did not stay long enough, 6 received postnatal home follow-up, 2 had to stay longer in hospital, 29 transferred to early discharge group due to bed shortages)
Selective reporting (re- porting bias)	Unclear risk	Not common practice to publish protocol at that time
Other bias	Low risk	Nothing to indicate any other source of bias

Taiba 2012

Study characteristics	
Methods	Design: RCT Setting: Dhaka, Bangladesh AN/PN recruitment: not reported
Participants	Number of women eligible: 300 Number of participants randomised to treatment groups: A: 150; B: 150. Number of post-randomisation exclusions: 0 Number of women who remained in trial (per group if available): A) B) Number of withdrawals (per group if available): A: 100/150; B: 0/150. Number of women analysed (per group if available): A: 50; B: 150. Gestational age at delivery (weeks): not reported Inclusion criteria: term pregnant women undergoing elective or emergency caesarean section Exclusion criteria: not reported
Interventions	A: (n = 150) discharge at 72 hours after caesarean B: (n = 150) discharge at 7 days after caesarean Duration of follow-up: 7 days after caesarean.
Outcomes	Maternal: Urinary and respiratory tract infections wound infection puerperal sepsis Infant: Umbilical cord sepsis



Taiba 2012 (Continued)

Septicemia respiratory distress syndrome

Notes	Co-interventions: not reported
	Dates of study: July 2006 to December 2006
	Funding sources: Not reported
	Declarations of interest: Not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	"Randomized" used in title but no description of randomisation."300 consecu- tive cases fulfilling the enrolment criteria were included in the study"
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible but lack of blinding not likely to have an effect on the outcomes measured in the trial
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	100/150 withdrawals in early discharge group, no withdrawals in 7 day group.
Selective reporting (re- porting bias)	Unclear risk	No protocol available. No indication of what the pre-specified outcomes were.
Other bias	Low risk	No indication of any other source of bias

Thompson 1999

Study characteristics

Methods	Design: RCT Setting: Canberra Hospital, Australia
	AN recruitment
Participants	Number of women eligible: 118
	Number of participants randomised to treatment groups: 50 (A: 24; B: 26)
	Number of post-randomisation exclusions: A: 13; B: 5 excluded postpartum and 3 antenatal with- drawals but no explanation for these 3 withdrawals.
	Number of women who remained in trial (per group if available): A: 11 (remained eligible for early dis- charge); B: 18 (remained eligible for early discharge)
	Number of withdrawals (described in trial as "did not comply"): A: 8/11; B: 9/18
	Number of women analysed (per group if available): A: 11; B: 18.
	Inclusion criteria: "women eligible for early discharge (i.e. not classified as high risk and not having a planned caesarean delivery)"
	Exclusion criteria: "Intrapartum exclusion criteria included caesarean delivery or postpartum haemon rhage.



Thompson 1999 (Continued)

Postpartum exclusion criteria for the mother included inability to care for herself or her infant or any medical conditions requiring close medical supervision. Postpartum exclusion criteria for the baby included jaundice requiring phototherapy, admission to the special care or neonatal intensive care unit, or both.

Interventions	A: (n = 24) length of stay 36 hours or less B: (n = 26) length of stay 37-72 hours Duration of follow-up: 72 hours after giving birth
Outcomes	Women's satisfaction with group allocation
Notes	Co-interventions: not reported Dates of study: "a six week period in 1996" Funding sources: "This project was supported from donations by the Salaried Special" Declarations of interest: not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"Women were randomly allocated to groups 1 or 2 at the time of recruitment. The allocation sequence was generated from random number table"
Allocation concealment (selection bias)	Low risk	Allocation was "concealed in sequentially numbered, opaque sealed envelopes"
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Blinding not possible. lack of blinding is likely to have an effect on the out- comes measured in the trial.
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	No clinical outcomes were measured
Incomplete outcome data (attrition bias) All outcomes	Low risk	13/24 and 5/26 postpartum exclusions (i.e. no longer eligible for early dis- charge) 3/11 and 9/18 complied. No clinical outcomes were measured so un- likely to be any risk of attrition bias
Selective reporting (re- porting bias)	High risk	The trial was designed to test a null hypothesis of no significant difference be- tween the frequency of postnatal depression in mothers discharged from hos- pital within 6 hours (A) compared with those discharged 37-72 hours after de- livery (B). This outcome was not reported.
Other bias	Low risk	Nothing to indicate any other source of bias

Waldenström 1987

Study characteristi	cs
Methods	Method of randomisation: unclear. AN recruitment
	Blinding: women and caregivers not blinded; blinding of outcome assessment - unclear.
	Follow-up: women to 6 weeks; infants to 6 months.



Waldenström 1987 (Continued)	Loss to follow-up: 60/164 (36.6%); 47 post-randomisation exclusions; 13 withdrawals in early discharge group; 100% response to 6-week questionnaires.
	Analysis: not intention to treat.
	Duration: March 1984 to June 1985.
	Setting: teaching hospital, Uppsala, Sweden.
Participants	Number of women eligible: 1604 Number of participants randomised to treatment groups: 164 (A: 85; B: 79) Number of post-randomisation exclusions: A: 26% (22?); B: 32% (25?) Number of women who remained in trial: A: 63; B: 54. Number of withdrawals:
	A: 13 (various reasons for choosing to stay in hospital longer)
	B: 0 (5 left hospital early but were included in analysis as per allocated group) Number of women analysed: A: 50; B: 54.
	Inclusion criteria: pregnancy and birth free from significant complications; vaginal delivery; singleton; gestational age > 37 weeks, birthweight >= 3000 g, Apgar >= 7 at 5 min; and no significant infant or ma- ternal morbidity in first 24 hours. Characteristics of participants: mean maternal age: early 28, standard 27. Proportion primiparous: early 20%, standard 30%. Maternal university education: early 28%, standard 19%. Mean birthweight: early 3658 g, standard 3481 g.
	In comparison with non-participants, trial participants had less education, were more 'family-oriented' and confident about parenthood, and more negative about care in hospital.
Interventions	A: (n = 85) discharge 24 to 48 hours. B: (n = 79) discharge 6 days after giving birth
	Co-interventions: Early group - nurse home visit 4 weeks before term; visit to hospital on day 5 for paediatric examina- tion; daily nurse home visits for 3 to 4 days post discharge. Standard group: traditional hospital care and no home visits post-discharge.
	Duration of follow-up: infants followed up for 6 months, women followed up for 6 weeks.
Outcomes	Infant and maternal re-admissions within 6 and 8 weeks pp respectively. Maternal depressed mood in first 6 weeks. Breastfeeding at 2 and 6 months. Maternal fatigue in first 14 days. Maternal satisfaction with care.
Notes	Mean length of stay at time of study was 6 days; but shorter in standard discharge group where mean stay was 4.1 nights. Women who 'crossed over' were treated differently in early and standard groups: 13 women excluded because they went home later than allocation, whereas 5 women allocated to standard discharge who went home early (but without home visits), were retained in analysis.
	Dates of study: March 1984 to June 1985
	Funding sources: "This study was supported by grants from The Swedish Ministry of Health and Social Affairs, Commission for Social Research (Project no. F 83/18)"
	Declarations of interest: not reported
Risk of bias	



Waldenström 1987 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Insufficient information in the report to assess whether sequence generation was adequate.
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcome of hos- pital readmission
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Unclear for outcome assessors.
Incomplete outcome data	High risk	More missing data from early discharge group than control group.
(attrition bias) All outcomes		13/63 women in the early discharge group chose to stay longer in hospital) and were not included in analysis, except for data obtained from Child Health Center records pertaining to infant morbidity and breastfeeding).
		5/54 in the control group chose to leave hospital early and were included in analysis.
Selective reporting (re- porting bias)	Unclear risk	Not common practice to publish protocol at that time
Other bias	Low risk	Nothing to indicate any other source of bias

Winterburn 2000

Method of randomisation: via "selecting sealed envelopes"; process unclear. Antenatal recruitment
Blinding: women and caregivers not blinded; unclear for outcome assessment.
Analysis: by intention to treat and on basis of actual length of stay.
Loss to follow-up: 7/255 (2.7%).
Follow-up: to 1 month pp.
Duration: February 1996 to June 1998.
Setting: large teaching hospital, north of England.
Number of women eligible: not reported Number of participants randomised to treatment groups: 248 (A: 121; B: 127) Number of post-randomisation exclusions: 0 Number of women who remained in trial: A: 121; B: 127. Number of withdrawals: 7 withdrew for reasons not related to the intervention (study does not report which groups they were in) Number of women analysed: A: 121; B: 127.



Winterburn 2000 (Continued)

Trusted evidence. Informed decisions. Better health.

Ninterburn 2000 (Continued)		time mothers wanting to breastfeed and with no preference about length of hos- early discharge criteria.	
	Characteristics of parti no information about s	icipants: socio-demographic characteristics.	
Interventions	A: (n = 121) hospital sta B: (n = 127) hospital sta		
	Co-interventions: community midwife ho not specified).	ome visits to support breastfeeding (number of visits and over what time period	
Outcomes	Proportion of women b	preastfeeding.	
Notes	Major limitation due to cross-over of study participants - in both directions, resulting in only 51 women experiencing early discharge and 197 experiencing standard discharge. Unclear whether home visits offered to all women who went home < 48 hours, regardless of allocated group status.		
	Dates of study: February 1996 to June 1998		
	Funding sources: "The study was supported by a grant from the Northern General Hospital Trust Re- search Committee."		
	Declarations of interest: not reported		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Unclear risk	Insufficient information in the report to assess whether sequence generation was adequate.	
Allocation concealment (selection bias)	Unclear risk	Not reported	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcomes mea- sured.	
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Unclear for outcome assessors.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	7/155 women withdrew for reasons not related to duration of hospital stay.All other women analysed according to allocated group.	
Selective reporting (re- porting bias)	Unclear risk	Not common practice to publish protocol at that time	



Yanover 1976

Random sequence genera- tion (selection bias)	Unclear risk	Insufficient information in the report to assess whether sequence generation was adequate.	
Bias	Authors' judgement	Support for judgement	
Risk of bias			
	Declarations of interest: Not reported		
	Funding sources: "Supported by Kaiser Foundation Research Institute, General Research Support, un- der a grant (5 501 RR05521-10) from the US Public Health Service		
	Dates of study: Not reported		
Notes	Highly selected study p	articipants; cross-over a problem.	
Outcomes	Infant and maternal re-admissions to 6 weeks pp. Maternal views about length of hospital stay.		
	Duration of follow-up: 6	5 weeks	
	Co-interventions: A: nursing staff intensively trained for early discharge; prospective parents in early group attended pre- natal early discharge preparation classes; daily home visits through 4th day pp. B: received prenatal education; paediatric visit at 2 weeks pp; and an obstetric visit at 6 weeks.		
Interventions	hours, range = 12-86 ho	m 12 to 48 hours pp (12 were discharged later than 48 hours); median stay 26 urs. > 48 hours pp (5 discharged at < 48 hours); median stay = 68 hours, range = 31 t	
	ration of marriage, time	cipants: groups on maternal age, race, father's occupation, planned pregnancy du- e to conceive, maternal and paternal education, presence of another child at ences for infant feeding, prenatal education or natural childbirth; BUT no data	
	father willing to attend	/ 0 or 1; maternal age 19 to 35; low medical risk; at least 12th grade education; prenatal classes; prospective parents living together; adequate English; living l; and assessment of mother and infant as eligible for early discharge (range of	
Participants	Number of post-randor premature labour, 2 cae Number of women who	randomised to treatment groups: 128 randomised nisation exclusions (not reported per group): 15 (4 stillbirth, 4 pre-eclampsia, 2 esarean sections, 3 'other') remained in trial: A: 44 B: 44 ; (not reported per group): 25	
	Setting: Departments o Francisco, USA.	f Obstetrics, Pediatrics and Nursing, Kaiser-Permanente Medical Center, San	
	Antenatal recruitment		
Methods	RCT		



Yanover 1976 (Continued)

Allocation concealment (selection bias)	Unclear risk	Not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Blinding not possible for mothers but unlikely to influence the outcome of hos- pital readmissions
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Unclear for outcome assessors.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	15/128 excluded for no longer meeting early discharge criteria, and 25/128 withdrew due to reasons unrelated to duration of hospital stay but the study authors do not report the numbers of exclusions and withdrawals per group.
Selective reporting (re- porting bias)	Unclear risk	Not common practice to publish protocol at that time
Other bias	Low risk	Nothing to indicate any other source of bias

C: control CHF: Swiss francs CS: caesarean section hrs: hours I: intervention min: minutes pp: postpartum RCT: randomised controlled trial SD: standard deviation

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Escobar 2001	Women participating in this trial were randomised to home visits or hospital-based group fol- low-up visits after early obstetric discharge, i.e. randomisation was not used to compare early with longer length of stay.
Lieu 2000	Women participating in this trial were randomised to home visits or hospital-based group fol- low-up visits after early obstetric discharge, i.e. randomisation was not used to compare early with longer length of stay.
Steel O'Connor 2003	Women participating in this trial were randomised to public health nurse follow-up or a screening telephone call after early obstetric discharge, i.e. randomisation was not used to compare early with longer length of stay.

Characteristics of studies awaiting classification [ordered by study ID]

NCT04422041 2020

Methods

Randomised controlled trial

Setting: Universidad Autonoma de Nuevo Leon, School of Medicine, Mexico



NCT04422041 2020 (Continued)	
Participants	Recruitment target: 354
	Inclusion criteria: healthy newborns that were born from vaginal delivery in primiparous or mul- tiparous women where both the mother and the newborn were deemed as eligible for early dis- charge according to the American Association of Pediatrics criteria and by a clinical obstetric moth- er evaluation.
	Exclusion criteria: placenta praevia, abnormal bleeding during vaginal delivery (considered as greater than 500mL), inability to deambulate, medical complications from previous a previous pregnancy, 3rd or 4th degree perineal laceration as well as medical conditions that required any monitorization for more than 24 hours after delivery.
Interventions	Early discharge: Time to discharge less than 24 hours after birth
	Comparator: Discharge time between 24 and 48 hours after birth
Outcomes	Readmission to hospital: proportion of infants readmitted within 28 days
	Proportion of newborns who attended the emergency services within 28 days
Notes	Registered June 2020.
	Trial reported as having taken place from July 2016 to June 2018.
	No published results available.

Characteristics of ongoing studies [ordered by study ID]

Namagembe 2014

Maternal outcome following discharge on 2nd versus 3rd day post caesarean section at Mulago Hospital
Parallel, 2 arm RCT.
Setting: Kampala, Uganda
Mothers who had uncomplicated Caesarean section, will be randomised after 24 hours after the operation and those who fall on the control group will be discharged after 72 hours
Recruitment target: 338
Inclusion criteria:
 Women age 18-45, within 20km of Mulago Postnatal mothers who will consent to participate in the study
Exclusion criteria:
 Anybody with the condition which is likely to prolong her stay in Hospital beyond 3 days such as uncontrolled hypertension, diabetes mellitus, anaemia Hb < 10g/dl, antepartum haemorrhages, premature rupture of membranes, obstructed labour
Suspected bladder injury, postpartum haemorrhage.
 Evidence of infection at the time of discharge (heart rate > 110 beats/min, temperature > 37.5oC) No telephone contact number for follow-up
Group 1: day 2 discharge
Group 2: day 3 discharge

Namagembe 2014 (Continued)

Outcomes	Primary outcome: wound infection
	Secondary outcomes: postpartum haemorrhage, wound dehiscence, acceptability with the hospi- tal discharge
Starting date	January 2014
Contact information	Cecilia Thomas cecyliamasili@yahoo.com
	Imelda Namagembe namagime@gmail.com
Notes	Funding source: St Augustine University of Tanzania
	Study authors were contacted by email but the study authors did not reply

DATA AND ANALYSES

Comparison 1. Early versus standard discharge

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1.1 Infants readmitted for neonatal morbidity within 7 days	2	247	Risk Ratio (M-H, Fixed, 95% CI)	1.15 [0.42, 3.16]
1.2 Infants readmitted for neonatal morbidity within 28 days	10	6918	Risk Ratio (M-H, Fixed, 95% CI)	1.59 [1.27, 1.98]
1.2.1 Trials conducted in 1960s	1	2151	Risk Ratio (M-H, Fixed, 95% CI)	1.83 [0.43, 7.80]
1.2.2 Trials conducted in 1970s	1	88	Risk Ratio (M-H, Fixed, 95% CI)	5.00 [0.25, 101.25]
1.2.3 Trials conducted in 1980s	2	226	Risk Ratio (M-H, Fixed, 95% CI)	0.44 [0.12, 1.63]
1.2.4 Trials conducted in 1990s	2	540	Risk Ratio (M-H, Fixed, 95% CI)	2.43 [0.87, 6.79]
1.2.5 Trials conducted in 2000s	1	430	Risk Ratio (M-H, Fixed, 95% CI)	0.61 [0.15, 2.53]
1.2.6 Trials conducted in 2010s	3	3483	Risk Ratio (M-H, Fixed, 95% Cl)	1.65 [1.30, 2.10]
1.3 Infants readmitted for neonatal morbidity within 28 days: mode of birth subgroups	10	6918	Risk Ratio (M-H, Fixed, 95% CI)	1.59 [1.27, 1.98]
1.3.1 Vaginal birth	5	2854	Risk Ratio (M-H, Fixed, 95% CI)	1.30 [0.55, 3.09]



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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1.3.2 Caesarean birth	4	3605	Risk Ratio (M-H, Fixed, 95% CI)	1.57 [1.24, 1.99]
1.3.3 Vaginal or caesarean	1	459	Risk Ratio (M-H, Fixed, 95% CI)	2.43 [0.87, 6.79]
1.4 Infants readmitted for neonatal morbidity within 28 days: subgroups < 24h vs > 24h	10	6918	Risk Ratio (M-H, Fixed, 95% CI)	1.59 [1.27, 1.98]
1.4.1 Early discharge < 24 hours	3	3770	Risk Ratio (M-H, Fixed, 95% CI)	1.64 [1.28, 2.09]
1.4.2 Early discharge > 24 hours	6	3060	Risk Ratio (M-H, Fixed, 95% CI)	1.28 [0.73, 2.25]
1.4.3 Early discharge range < 24 hours and > 24 hours	1	88	Risk Ratio (M-H, Fixed, 95% Cl)	5.00 [0.25, 101.25]
1.5 Women readmitted within six weeks	11	6992	Risk Ratio (M-H, Fixed, 95% CI)	1.12 [0.82, 1.54]
1.5.1 Trials conducted in 1960s	1	2094	Risk Ratio (M-H, Fixed, 95% CI)	2.84 [0.68, 11.81]
1.5.2 Trials conducted in 1970s	1	88	Risk Ratio (M-H, Fixed, 95% CI)	Not estimable
1.5.3 Trials conducted in 1980s	3	357	Risk Ratio (M-H, Fixed, 95% CI)	0.25 [0.05, 1.36]
1.5.4 Trials conducted in 1990s	2	540	Risk Ratio (M-H, Fixed, 95% CI)	2.03 [0.37, 10.95]
1.5.5 Trials conducted in 2000s	1	430	Risk Ratio (M-H, Fixed, 95% Cl)	0.82 [0.22, 2.99]
1.5.6 Trials conducted in 2010s	3	3483	Risk Ratio (M-H, Fixed, 95% CI)	1.11 [0.78, 1.58]
1.6 Women readmitted within six weeks: mode of birth subgroups	11	6992	Risk Ratio (M-H, Fixed, 95% Cl)	1.12 [0.82, 1.54]
1.6.1 Vaginal birth	6	2928	Risk Ratio (M-H, Fixed, 95% CI)	1.32 [0.58, 3.02]
1.6.2 Caesarean section	4	3605	Risk Ratio (M-H, Fixed, 95% CI)	1.05 [0.74, 1.49]
1.6.3 Vaginal and caesarean	1	459	Risk Ratio (M-H, Fixed, 95% CI)	2.03 [0.37, 10.95]
1.7 Women readmitted within six weeks: subgroups < 24 hours vs > 24hrs	11	6992	Risk Ratio (M-H, Fixed, 95% CI)	1.12 [0.82, 1.54]



Cochrane Database of Systematic Reviews

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1.7.1 Early discharge < 24 hours	4	3858	Risk Ratio (M-H, Fixed, 95% CI)	1.08 [0.76, 1.54]
1.7.2 Early discharge > 24 hours	7	3134	Risk Ratio (M-H, Fixed, 95% CI)	1.28 [0.65, 2.55]
1.8 Women probably depressed within six months	5	4333	Risk Ratio (M-H, Random, 95% CI)	0.80 [0.46, 1.42]
1.8.1 Trials conducted in 1980s	1	104	Risk Ratio (M-H, Random, 95% CI)	0.65 [0.16, 2.57]
1.8.2 Trials conducted in 1990s	1	459	Risk Ratio (M-H, Random, 95% CI)	0.77 [0.41, 1.44]
1.8.3 Trials conducted in 2000s	1	430	Risk Ratio (M-H, Random, 95% CI)	0.25 [0.05, 1.19]
1.8.4 Trials conducted in 2010s	2	3340	Risk Ratio (M-H, Random, 95% CI)	1.08 [0.44, 2.64]
1.9 Women probably depressed within six months: mode of birth subgroups	5	4333	Risk Ratio (M-H, Random, 95% CI)	0.80 [0.46, 1.42]
1.9.1 Vaginal birth	2	534	Risk Ratio (M-H, Random, 95% CI)	0.43 [0.15, 1.19]
1.9.2 Caesarean birth	2	3340	Risk Ratio (M-H, Random, 95% CI)	1.08 [0.44, 2.64]
1.9.3 Vaginal or caesarean birth	1	459	Risk Ratio (M-H, Random, 95% CI)	0.77 [0.41, 1.44]
1.10 Women probably depressed with- in six months: subgroups < 24 h vs < 24 hrs	5	4333	Risk Ratio (M-H, Random, 95% CI)	0.80 [0.46, 1.42]
1.10.1 Early discharge < 24 hours	3	3770	Risk Ratio (M-H, Random, 95% CI)	0.62 [0.18, 2.16]
1.10.2 Early discharge > 24 hours	2	563	Risk Ratio (M-H, Random, 95% CI)	0.75 [0.42, 1.32]
1.11 Women breastfeeding (exclusively or partially) at six weeks postpartum	10	7156	Risk Ratio (M-H, Random, 95% CI)	1.04 [0.96, 1.13]
1.11.1 Trials conducted in 1960s	1	2257	Risk Ratio (M-H, Random, 95% CI)	2.49 [1.59, 3.91]
1.11.2 Trials conducted in 1980s	2	198	Risk Ratio (M-H, Random, 95% CI)	0.97 [0.78, 1.21]
1.11.3 Trials conducted in 1990s	4	931	Risk Ratio (M-H, Random, 95% CI)	1.04 [0.90, 1.19]



Cochrane Database of Systematic Reviews

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size	
1.11.4 Trials conducted in 2000s	1	430	Risk Ratio (M-H, Random, 95% CI)	1.06 [0.99, 1.15]	
1.11.5 Trials conducted in 2010s	2	3340	Risk Ratio (M-H, Random, 95% CI)	0.99 [0.83, 1.18]	
1.12 Women breastfeeding (exclusively or partially) at six weeks postpartum: mode of birth subgroups	10	7156	Risk Ratio (M-H, Random, 95% CI)	1.04 [0.96, 1.13]	
1.12.1 Vaginal birth	6	3112	Risk Ratio (M-H, Random, 95% CI)	1.15 [0.90, 1.47]	
1.12.2 Caesarean birth	2	3340	Risk Ratio (M-H, Random, 95% CI)	0.99 [0.83, 1.18]	
1.12.3 Vaginal or caesarean birth or not specified	2	704	Risk Ratio (M-H, Random, 95% CI)	1.03 [0.95, 1.11]	
1.13 Women breastfeeding (exclusively or partially) at six weeks postpartum: subgroups < 24hr vs > 24 hrs	10	7156	Risk Ratio (M-H, Random, 95% CI)	1.04 [0.96, 1.13]	
.13.1 Early discharge < 24 hours	3	3770	Risk Ratio (M-H, Random, 95% CI)	1.01 [0.90, 1.15]	
1.13.2 Early discharge > 24 hours	4	2866	Risk Ratio (M-H, Random, 95% CI)	1.12 [0.77, 1.63]	
1.13.3 Early discharge range < 24 hours and > 24 hours	3	520	Risk Ratio (M-H, Random, 95% CI)	1.10 [0.90, 1.34]	
1.14 Women breastfeeding (exclusively or partially) at 12 weeks postpartum	1	430	Risk Ratio (M-H, Fixed, 95% CI)	1.21 [1.03, 1.41]	
1.15 Women breastfeeding (partially or exclusively) at six months postpartum	3	973	Risk Ratio (M-H, Random, 95% CI)	1.11 [0.87, 1.43]	
1.16 Infant mortality within 28 days	2	4882	Risk Ratio (M-H, Fixed, 95% CI)	0.39 [0.04, 3.74]	
1.17 Infant mortality within one year	2	1986	Risk Ratio (M-H, Fixed, 95% CI)	0.45 [0.07, 2.77]	
1.18 Number of contacts with health- care professionals regarding infant health issues within four weeks of birth	3	639	Risk Ratio (M-H, Fixed, 95% CI)	0.88 [0.67, 1.16]	
19 Number of contacts with health- care professionals regarding maternal nealth issues within six weeks of birth	2	464	Risk Ratio (M-H, Fixed, 95% CI)	0.72 [0.43, 1.20]	
20 Women reporting health prob- ems (including perineal pain, perineal nfection, breast soreness, breast infec- ion, caesarean wound pain, caesarean	1	200	Risk Ratio (M-H, Fixed, 95% CI)	0.25 [0.11, 0.59]	



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
wound infection) in the first six weeks postpartum				
1.21 Women reporting infant feeding problems	2	2405	Risk Ratio (M-H, Random, 95% CI)	0.89 [0.43, 1.86]
1.22 Women satisfied with postnatal care - dichotomous data	4	3098	Risk Ratio (M-H, Random, 95% CI)	1.10 [0.95, 1.29]
1.23 Satisfaction with postnatal care - continuous data	2	306	Std. Mean Difference (IV, Fixed, 95% CI)	0.74 [0.50, 0.98]
1.24 Women who perceive their length of hospital stay as too short	1	82	Risk Ratio (M-H, Fixed, 95% CI)	2.00 [0.19, 21.21]
1.25 Women perceive their length of hospital stay as too long	1	82	Risk Ratio (M-H, Fixed, 95% CI)	0.56 [0.20, 1.52]

Analysis 1.1. Comparison 1: Early versus standard discharge, Outcome 1: Infants readmitted for neonatal morbidity within 7 days

	Early dis	scharge	Standard d	ischarge		Risk Ratio	Risk F	latio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed	l, 95% CI
Waldenström 1987	1	50	0	54	7.4%	3.24 [0.13 , 77.63]		
Kruse 2021	6	72	6	71	92.6%	0.99 [0.33 , 2.91]		F
Total (95% CI)		122		125	100.0%	1.15 [0.42 , 3.16]		
Total events:	7		6					
Heterogeneity: Chi ² = 0).48, df = 1 (I	P = 0.49); I	$^{2} = 0\%$			0.0	01 0.1 1	10 100
Test for overall effect: 2	Z = 0.28 (P =	0.78)				Favours	early discharge	Favours standard discharg
Test for subgroup differ	rences: Not a	pplicable						

14 20 20 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	all Eve 818 8 818 44 44 44 50 61 111 111	2 2 0 0 0	Total 333 333 44 44	2.9% 2.9% 0.4% 0.4%	M-H, Fixed, 95% CI 1.83 [0.43 , 7.80] 1.83 [0.43 , 7.80] 5.00 [0.25 , 101.25] 5.00 [0.25 , 101.25]	M-H, Fixed, 95% CI	A B C D E F ● ? ● ? ? ? ? → ? ? ● ? ? ? ?
14 20 2 (0.41) 2 2 (0.29) 1 2 3 P = 0.47);	44 44 44 50 61	2 0 0	333 44 44	2.9% 0.4%	1.83 [0.43 , 7.80] 5.00 [0.25 , 101.25]		
14 20 2 (0.41) 2 2 (0.29) 1 2 3 P = 0.47);	44 44 44 50 61	2 0 0	333 44 44	2.9% 0.4%	1.83 [0.43 , 7.80] 5.00 [0.25 , 101.25]		
20 2 0.41) 2 2 2 0.29) 1 2 3 P = 0.47);	44 44 50 61	0 0	44 44	0.4%	5.00 [0.25 , 101.25]		→ 22•22 -
2 0.41) 2 2 2 0.29) 1 2 3 P = 0.47);	44 50 61	0 0	44				→
2 2 0.29) 1 2 3 P = 0.47);	44 50 61	0	44				→ ?? ? •??? -
2 2 0.29) 1 2 3 P = 0.47);	44 50 61	0	44				→ ?? ? •??? -
2 : 0.29) 1 2 3 P = 0.47);	44 50 61	0	44				→ ?? ? ??? -
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2 : 0.29) 1 2 3 P = 0.47);	44 50 61	0	44				-
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1 2 3 P = 0.47);	61	1					
1 2 3 P = 0.47);	61						
2 3 P = 0.47);	61		<i></i>				
2 3 P = 0.47);	61						
2 3 P = 0.47);	61		-				
3 P = 0.47);		6	54	0.8%	1.08 [0.07 , 16.81]		? 🗧 ? 🖶 ? 🔵 ?
3 P = 0.47);	111		61	5.1%	0.33 [0.07 , 1.59]		🖶 🖶 🖶 ? ???
P = 0.47);			115	5.9%	0.44 [0.12 , 1.63]		
		7				-	
0.22)	; I ² = 0%						
,							
0	35	0	46		Not estimable		?? 🕂 ? 🕂 ?
12 2	228	5	231	4.2%	2.43 [0.87 , 6.79]		+++?+
2	263		277	4.2%	2.43 [0.87 , 6.79]		
12		5				-	
0.09)							
3	213	5	217	4 2%	0.61 [0.15 2 53]		? 🗭 🖨 ? ? ?
		5					•••••
	_10	5	21/	/0	0.01 [0.10 , 2.00]		
J		э					
0.50)							
3	170	4	170	3 /0/	0.76[0.17 3.34]		
		б					
	137		1746	82.5%	1.65 [1.30 , 2.10]	•	
	**	98					
	; 1² = 4%						
A [*]	186		2732	100.0%	1.59 [1.27 - 1.98]		
		117	27.52	10000/0	100 [1127 , 1100]	▼	
	I2 - 100/	11/			F		
	, 1- = 12%						100
() () ()	3 3 52 1 6 1 61 P = 0.35) < 0.0001) 4 01 P = 0.33) < 0.0001)	213 3 (3, 50) (3, 170) (52, 1495) (6, 72) (1737) (61) $P = 0.35); 1^2 = 4\%$ (3, 0001) (12)	213 3 5 (3, 5) (3, 170) 4 (52, 1495) 88 (6, 72) 6 (72, 6) (737) 98 (2, 0, 001) 98 (2, 173) 98 (2, 173) 98 (3,	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	213 217 4.2% 3 5 $a,50$ 3 170 4 172 3.4% 52 1495 88 1503 74.0% 6 72 6 71 1737 1746 82.5% 61 98 P = 0.35); I ² = 4% $a,0001$ 117 P = 0.33); I ² = 12% $a,0001$	213 217 4.2% 0.61 [0.15, 2.53] 3 5 $= 0.50$) 3 170 4 172 3.4% 0.76 [0.17, 3.34] $= 0.50$) 3 170 4 172 3.4% 0.76 [0.17, 3.34] $= 0.50$) 3 170 4 172 3.4% 0.76 [0.17, 3.34] 52 1495 88 1503 74.0% 1.74 [1.35, 2.24] 6 72 6 71 5.1% 0.99 [0.33, 2.91] 1737 1746 82.5% 1.65 [1.30, 2.10] 61 98 98 98 98 $P = 0.355$); $I^2 = 4\%$ 2732 100.0% 1.59 [1.27, 1.98] 01 117 117 0.01 $P = 0.333$); $I^2 = 12\%$ 0.01 5732 100.0% 1.59 [1.27, 1.98] 01 117 117 0.01 54000000000000000000000000000000000000	213 3 5 3 5 (-50) 3 170 4 172 3.4% 0.76 [0.17, 3.34] 52 1495 88 1503 74.0% 1.74 [1.35, 2.24] 6 72 6 71 5.1% 0.99 [0.33, 2.91] 1737 1746 82.5% 1.65 [1.30, 2.10] 61 98 P = 0.35); I ² = 4% (-0,0001) 4186 2732 100.0% 1.59 [1.27, 1.98] 01 117 P = 0.33); I ² = 12% (-0,01 - 0,1 - 1 - 10) Favours early discharge Favours stand

Analysis 1.2. Comparison 1: Early versus standard discharge, Outcome 2: Infants readmitted for neonatal morbidity within 28 days

Footnotes

(1) Measured at six weeks postpartum

(2) Measured at seven days postpartum

(3) Measured at one month postpartum

(4) Measured at eight weeks postpartum

Risk of bias legend

(A) Random sequence generation (selection bias)

(B) Allocation concealment (selection bias)

(C) Blinding of participants and personnel (performance bias)

(D) Blinding of outcome assessment (detection bias)

(E) Incomplete outcome data (attrition bias): All outcomes



Analysis 1.2. (Continued)

(D) Blinding of outcome assessment (detection bias)(E) Incomplete outcome data (attrition bias): All outcomes(F) Selective reporting (reporting bias)

(G) Other bias

Analysis 1.3. Comparison 1: Early versus standard discharge, Outcome 3: Infants readmitted for neonatal morbidity within 28 days: mode of birth subgroups

	Early dis	scharge	Standard d	ischarge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
1.3.1 Vaginal birth							
Hellman 1962	20	1818	2	333	2.9%	1.83 [0.43 , 7.80]	_
Yanover 1976 (1)	2	44	0	44	0.4%	5.00 [0.25 , 101.25]	}
Waldenström 1987 (2)	1	50	1	54	0.8%	1.08 [0.07 , 16.81]	← →
Smith-Hanrahan 1995 (1)	0	35	0	46		Not estimable	
Sainz Bueno 2005	3	213	5	217	4.2%	0.61 [0.15 , 2.53]	←
Subtotal (95% CI)		2160		694	8.3%	1.30 [0.55 , 3.09]	
Total events:	26		8				
Heterogeneity: Chi ² = 2.09,	df = 3 (P =	0.55); I ² =	0%				
Test for overall effect: $Z = 0$	0.60 (P = 0.	55)					
1.3.2 Caesarean birth							
Brooten 1994	2	61	6	61	5.1%	0.33 [0.07 , 1.59]	← → −
Chiong 2012 (3)	3	170	4	172	3.4%	0.76 [0.17 , 3.34]	• • •
Bayoumi 2016 (1)	152	1495	88	1503	74.0%	1.74 [1.35 , 2.24]	
Kruse 2021	6	72	6	71	5.1%	0.99 [0.33 , 2.91]	
Subtotal (95% CI)		1798		1807	87.5%	1.57 [1.24 , 1.99]	•
Total events:	163		104				•
Heterogeneity: Chi ² = 6.03,	df = 3 (P =	0.11); I ² =	50%				
Test for overall effect: $Z = 2$	3.76 (P = 0.)	0002)					
1.3.3 Vaginal or caesarear	1						
Boulvain 2004 (4)	12	228	5	231	4.2%	2.43 [0.87 , 6.79]	
Subtotal (95% CI)		228		231	4.2%	2.43 [0.87 , 6.79]	
Total events:	12		5				
Heterogeneity: Not applica	ble						
Test for overall effect: Z =	1.70 (P = 0.	09)					
Total (95% CI)		4186		2732	100.0%	1.59 [1.27 , 1.98]	
Total events:	201		117				-
Heterogeneity: Chi ² = 9.10,	df = 8 (P =	0.33); I ² =	12%				
Test for overall effect: $Z = $	4.07 (P < 0.	0001)				Favou	rs early discharge Favours standard disc
est for subgroup differenc	es: $Chi^2 = 0$.87, df = 2	$(P = 0.65), I^2$	= 0%			

Footnotes

(1) Measured at six weeks postpartum

(2) Measured at seven days postpartum

(3) Measured at eight weeks postpartum

(4) Measured at one month postpartum

Analysis 1.4. Comparison 1: Early versus standard discharge, Outcome 4: Infants readmitted for neonatal morbidity within 28 days: subgroups < 24h vs > 24h

	Early dis	scharge	Standard d	ischarge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
1.4.1 Early discharge < 24	4 hours						
Sainz Bueno 2005	3	213	5	217	4.2%	0.61 [0.15 , 2.53]	_
Chiong 2012 (1)	3	170	4	172	3.4%	0.76 [0.17 , 3.34]	_
Bayoumi 2016 (2)	152	1495	88	1503	74.0%	1.74 [1.35 , 2.24]	
Subtotal (95% CI)		1878		1892	81.6%	1.64 [1.28 , 2.09]	▲
Total events:	158		97				•
Heterogeneity: Chi ² = 3.09,	, df = 2 (P =	0.21); I ² =	35%				
Test for overall effect: $Z = 2$	3.97 (P < 0.	0001)					
1.4.2 Early discharge > 24	1 hours						
Hellman 1962	20	1818	2	333	2.9%	1.83 [0.43 , 7.80]	
Waldenström 1987 (3)	1	50	1	54	0.8%	1.08 [0.07 , 16.81]	
Brooten 1994	2	61	6	61	5.1%	0.33 [0.07 , 1.59]	
Smith-Hanrahan 1995 (2)	0	35	0	46		Not estimable	
Boulvain 2004	12	228	5	231	4.2%	2.43 [0.87 , 6.79]	
Kruse 2021	6	72	6	71	5.1%	0.99 [0.33 , 2.91]	
Subtotal (95% CI)		2264		796	18.0%	1.28 [0.73 , 2.25]	
Total events:	41		20				
Heterogeneity: Chi ² = 4.83,	, df = 4 (P =	0.31); I ² =	17%				
Test for overall effect: Z =	0.85 (P = 0.	40)					
1.4.3 Early discharge ran	ge < 24 hou	rs and > 2	4 hours				
Yanover 1976 (2)	2	44	0	44	0.4%	5.00 [0.25 , 101.25]	
Subtotal (95% CI)		44		44	0.4%	5.00 [0.25 , 101.25]	
Total events:	2		0				
Heterogeneity: Not applica	ble						
Test for overall effect: Z =	1.05 (P = 0.	29)					
Total (95% CI)		4186		2732	100.0%	1.59 [1.27 , 1.98]	•
Total events:	201		117				•
Heterogeneity: Chi ² = 9.10,	, df = 8 (P =	0.33); I ² =	12%				0.01 0.1 1 10 100
Test for overall effect: Z =	4.07 (P < 0.	0001)				Favo	urs early discharge Favours standard dis
Test for subgroup differenc	CL 12 1	10 10 0	~ ~ ~ ~ ~				

Footnotes

(1) Measured at eight weeks postpartum

(2) Measured at six weeks postpartum

(3) Measured at seven days postpartum

Analysis 1.5. Comparison 1: Early versus standard discharge, Outcome 5: Women readmitted within six weeks

Study or Subgroup	Early dis Events	charge Total	Standard d Events	ischarge Total	Weight	Risk Ratio M-H, Fixed, 95% CI	Risk Ratio M-H, Fixed, 95% CI
	060-				-		
.5.1 Trials conducted in 19		1770	2	210	4 70/	2.04[0.00 11.01]	
Iellman 1962	32	1778	2	316	4.7%		+
ubtotal (95% CI)	22	1778	2	316	4.7%	2.84 [0.68 , 11.81]	
otal events:	32		2				
leterogeneity: Not applicab est for overall effect: Z = 1		15)					
.5.2 Trials conducted in 19	970s						
anover 1976	0	44	0	44		Not estimable	
ıbtotal (95% CI)		44		44		Not estimable	
otal events:	0		0				
eterogeneity: Not applicab	le						
est for overall effect: Not a	pplicable						
5.3 Trials conducted in 19	980s						
Valdenström 1987	0	50	1	54	2.0%		
arty 1990 (1)	1	93	1	38	2.0%		
rooten 1994	0	61	3	61	4.8%		
ubtotal (95% CI)		204		153	8.8%	0.25 [0.05 , 1.36]	
otal events:	1		5				-
eterogeneity: Chi ² = 0.31,	df = 2 (P =	0.86); I ² =	0%				
est for overall effect: $Z = 1$.61 (P = 0.	11)					
5.4 Trials conducted in 1							
mith-Hanrahan 1995	0	35	0	46		Not estimable	
oulvain 2004 (2)	4	228	2	231	2.7%		
ubtotal (95% CI)		263		277	2.7%	2.03 [0.37 , 10.95]	
otal events:	4		2				
eterogeneity: Not applicab							
est for overall effect: $Z = 0$.82 (P = 0.4	41)					
5.5 Trials conducted in 2		212	-	217	6.00/		
ainz Bueno 2005	4	213	5	217	6.8%	. , .	
ubtotal (95% CI)		213	-	217	6.8%	0.82 [0.22 , 2.99]	
otal events:	4		5				
teterogeneity: Not applicab est for overall effect: Z = 0		76)					
.5.6 Trials conducted in 20	010s						
hiong 2012	1	170	1	172	1.4%	1.01 [0.06 , 16.04]	
ayoumi 2016	56	1495	51	1503	70.1%		_
ruse 2021	5	72	4	71	5.5%		
ubtotal (95% CI)		1737		1746	77.0%		
otal events:	62		56	-			T
eterogeneity: $Chi^2 = 0.03$, est for overall effect: $Z = 0$	df = 2 (P =	-					
otal (95% CI)		4239		2753	100.0%	1.12 [0.82 , 1.54]	•
otal events:	103		70				ľ
eterogeneity: Chi ² = 5.27,	df = 8 (P =	0.73); I ² =	0%			0	1.001 0.1 1 10 100
est for overall effect: $Z = 0$							early discharge Favours standard disc
est for subgroup difference	`	· ·	$(P = 0.25)$ I^2	= 25 4%			

Footnotes

(1) Measured at 1 month postpartum(2) measured at 1 month postpartum



Analysis 1.6. Comparison 1: Early versus standard discharge, Outcome 6: Women readmitted within six weeks: mode of birth subgroups

	Early dis	charge	Standard d	ischarge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
1.6.1 Vaginal birth							
Hellman 1962	32	1778	2	316	4.7%	2.84 [0.68 , 11.81]	
Yanover 1976	0	44	0	44		Not estimable	
Waldenström 1987	0	50	1	54	2.0%	0.36 [0.01 , 8.63]	.
Carty 1990 (1)	1	93	1	38	2.0%	0.41 [0.03 , 6.37]	-
Smith-Hanrahan 1995	0	35	0	46		Not estimable	
Sainz Bueno 2005	4	213	5	217	6.8%	0.82 [0.22 , 2.99]	_
Subtotal (95% CI)		2213		715	15.4%	1.32 [0.58 , 3.02]	•
Total events:	37		9				-
Heterogeneity: Chi ² = 2.99	ə, df = 3 (P =	0.39); I ² =	0%				
Test for overall effect: Z =	0.65 (P = 0.	51)					
1.6.2 Caesarean section							
Brooten 1994	0	61	3	61	4.8%	0.14 [0.01 , 2.71]	_
Chiong 2012	1	170	1	172	1.4%	1.01 [0.06 , 16.04]	
Bayoumi 2016	56	1495	51	1503	70.1%	1.10 [0.76 , 1.60]	•
Kruse 2021	5	72	4	71	5.5%	1.23 [0.35 , 4.40]	
Subtotal (95% CI)		1798		1807	81.8%	1.05 [0.74 , 1.49]	•
Total events:	62		59				T
Heterogeneity: Chi ² = 1.89	ə, df = 3 (P =	0.60); I ² =	0%				
Test for overall effect: $Z =$	0.30 (P = 0.	77)					
1.6.3 Vaginal and caesare	ean						
Boulvain 2004 (2)	4	228	2	231	2.7%	2.03 [0.37 , 10.95]	_ _
Subtotal (95% CI)		228		231	2.7%	2.03 [0.37 , 10.95]	
Total events:	4		2				-
Heterogeneity: Not applica	able						
Test for overall effect: Z =	0.82 (P = 0.	41)					
Total (95% CI)		4239		2753	100.0%	1.12 [0.82 , 1.54]	•
Total events:	103		70				
Heterogeneity: Chi ² = 5.27	7, df = 8 (P =	0.73); I ² =	0%				-++++++ 0.01 0.1 1 10 100
Test for overall effect: Z =	0.72 (P = 0.	47)				Favou	rs early discharge Favours standard e

Test for subgroup differences: $Chi^2 = 0.74$, df = 2 (P = 0.69), $I^2 = 0\%$

Footnotes

(1) Measured at 1 month postpartum

(2) Measured at one month postpartum

Analysis 1.7. Comparison 1: Early versus standard discharge, Outcome 7: Women readmitted within six weeks: subgroups < 24 hours vs > 24hrs

	Early dis	scharge	Standard d	ischarge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
1.7.1 Early discharge < 2	24 hours						
Yanover 1976	0	44	0	44		Not estimable	
Sainz Bueno 2005	4	213	5	217	6.8%	0.82 [0.22 , 2.99]	
Chiong 2012	1	170	1	172	1.4%	1.01 [0.06 , 16.04]	
Bayoumi 2016	56	1495	51	1503	70.1%	1.10 [0.76 , 1.60]	_
Subtotal (95% CI)		1922		1936	78.3%	1.08 [0.76 , 1.54]	▲
Total events:	61		57				
Heterogeneity: Chi ² = 0.2	0, df = 2 (P =	0.91); I ² =	0%				
Test for overall effect: Z =	= 0.41 (P = 0.	68)					
1.7.2 Early discharge > 2	24 hours						
Hellman 1962	32	1778	2	316	4.7%	2.84 [0.68 , 11.81]	
Waldenström 1987	0	50	1	54	2.0%	0.36 [0.01 , 8.63]	.
Carty 1990 (1)	1	93	1	38	2.0%	0.41 [0.03 , 6.37]	
Brooten 1994	0	61	3	61	4.8%	0.14 [0.01 , 2.71]	
Smith-Hanrahan 1995	0	35	0	46		Not estimable	
Boulvain 2004 (2)	4	228	2	231	2.7%	2.03 [0.37 , 10.95]	
Kruse 2021	5	72	4	71	5.5%	1.23 [0.35 , 4.40]	_
Subtotal (95% CI)		2317		817	21.7%	1.28 [0.65 , 2.55]	•
Total events:	42		13				•
Heterogeneity: Chi ² = 4.9	1, df = 5 (P =	0.43); I ² =	0%				
Test for overall effect: Z =	= 0.71 (P = 0.	48)					
Total (95% CI)		4239		2753	100.0%	1.12 [0.82 , 1.54]	
Total events:	103		70				
Heterogeneity: Chi ² = 5.2	7, df = 8 (P =	0.73); I ² =	0%				0.01 0.1 1 10 100
Test for overall effect: Z =	= 0.72 (P = 0.	47)				Favou	rs early discharge Favours standard disc
Test for subgroup differen	ices: Chi ² = 0	.20, df = 1	$(P = 0.66), I^2 =$	= 0%			

Footnotes

(1) Measured at 1 month postpartum(2) measured at 1 month postpartum

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Analysis 1.8. Comparison 1: Early versus standard discharge, Outcome 8: Women probably depressed within six months

	Early dis	scharge	Standard d	ischarge		Risk Ratio	Risk Ratio	Risk of Bias
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI	ABCDEFG
.8.1 Trials conducted	in 1980s							
Waldenström 1987	3	50	5	54	12.2%	0.65 [0.16 , 2.57]		?? 🕂 ? 🛑 ? 🗧
ubtotal (95% CI)		50		54	12.2%	0.65 [0.16 , 2.57]		
otal events:	3		5					
eterogeneity: Not app	licable							
est for overall effect: 2	Z = 0.62 (P =	0.54)						
.8.2 Trials conducted	in 1990s							
300 Soulvain 2004	16	228	21	231	28.5%	0.77 [0.41 , 1.44]		+ + + ? + ? 4
ubtotal (95% CI)		228		231	28.5%	0.77 [0.41 , 1.44]	-	
otal events:	16		21					
leterogeneity: Not app	licable							
Test for overall effect: 2	Z = 0.81 (P =	0.42)						
.8.3 Trials conducted	in 2000s							
Sainz Bueno 2005	2	213	8	217	10.4%	0.25 [0.05 , 1.19]	_	? 🖶 🖶 ? ? ? 🖣
ubtotal (95% CI)		213		217	10.4%	0.25 [0.05 , 1.19]		
Total events:	2		8					
Heterogeneity: Not app	licable							
Cest for overall effect: 2	Z = 1.74 (P =	0.08)						
.8.4 Trials conducted	in 2010s							
Chiong 2012	1	170	3	172	5.5%	0.34 [0.04 , 3.21]	.	•••••••••••••••••••••••••••••••••••••••
Bayoumi 2016	1171	1495	914	1503	43.4%	1.29 [1.23 , 1.35]		? 🖶 🖶 🖶 ? 🧲
ubtotal (95% CI)		1665		1675	48.9%	1.08 [0.44 , 2.64]	-	
otal events:	1172		917				T	
Ieterogeneity: Tau ² = 0).24; Chi ² = 1	.36, df = 1	(P = 0.24); I ²	= 27%				
Test for overall effect: 2	Z = 0.16 (P =	0.87)						
Fotal (95% CI)		2156		2177	100.0%	0.80 [0.46 , 1.42]	•	
'otal events:	1193		951				· · · · · · · · · · · · · · · · · · ·	
Ieterogeneity: Tau ² = 0).19; Chi ² = 9	9.44, df = 4	(P = 0.05); I ²	= 58%			0.05 0.2 1 5 20	_
est for overall effect: 2	Z = 0.76 (P =	0.45)					Early discharge Standard disc	harge
est for subgroup differ	ences: Chi ² :	= 2.57. df =	3(P = 0.46).	$I^2 = 0\%$				

Risk of bias legend

(A) Random sequence generation (selection bias)

(B) Allocation concealment (selection bias)

(C) Blinding of participants and personnel (performance bias)

(D) Blinding of outcome assessment (detection bias)

(E) Incomplete outcome data (attrition bias): All outcomes

(F) Selective reporting (reporting bias)

(G) Other bias

Analysis 1.9. Comparison 1: Early versus standard discharge, Outcome 9: Women probably depressed within six months: mode of birth subgroups

	Early dis	charge	Standard d	ischarge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI
1.9.1 Vaginal birth							
Waldenström 1987	3	50	5	54	12.2%	0.65 [0.16 , 2.57]	
Sainz Bueno 2005	2	213	8	217	10.4%	0.25 [0.05 , 1.19]	_
Subtotal (95% CI)		263		271	22.5%	0.43 [0.15 , 1.19]	
Total events:	5		13				—
Heterogeneity: Tau ² = 0	.00; Chi ² = 0	.80, df = 1	(P = 0.37); I ²	= 0%			
Test for overall effect: Z	2 = 1.62 (P =	0.10)					
1.9.2 Caesarean birth							
Chiong 2012	1	170	3	172	5.5%	0.34 [0.04 , 3.21]	_
Bayoumi 2016	1171	1495	914	1503	43.4%	1.29 [1.23 , 1.35]	-
Subtotal (95% CI)		1665		1675	48.9%	1.08 [0.44 , 2.64]	
Total events:	1172		917				T
Heterogeneity: $Tau^2 = 0$.24; Chi ² = 1	.36, df = 1	(P = 0.24); I ²	= 27%			
Test for overall effect: Z	Z = 0.16 (P =	0.87)					
1.9.3 Vaginal or caesar	ean birth						
Boulvain 2004	16	228	21	231	28.5%	0.77 [0.41 , 1.44]	_ _ _
Subtotal (95% CI)		228		231	28.5%	0.77 [0.41 , 1.44]	
Total events:	16		21				~
Heterogeneity: Not app	licable						
Test for overall effect: Z	Z = 0.81 (P =	0.42)					
Total (95% CI)		2156		2177	100.0%	0.80 [0.46 , 1.42]	•
Total events:	1193		951				-
Heterogeneity: Tau ² = 0	.19; Chi ² = 9	.44, df = 4	(P = 0.05); I ²	= 58%			-++++++++++++++++++++++++++++++++++++
Test for overall effect: Z	z = 0.76 (P =	0.45)					Early discharge Standard dischar
Test for subgroup differ	ences: Chi² =	= 1.79, df =	2 (P = 0.41),	$I^2 = 0\%$			

Analysis 1.10. Comparison 1: Early versus standard discharge, Outcome 10: Women probably depressed within six months: subgroups < 24 h vs < 24 hrs

	Early dis	charge	Standard di	ischarge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI
1.10.1 Early discharge	< 24 hours						
Sainz Bueno 2005	2	213	8	217	10.4%	0.25 [0.05 , 1.19]	_
Chiong 2012	1	170	3	172	5.5%	0.34 [0.04 , 3.21]	
Bayoumi 2016	1171	1495	914	1503	43.4%	1.29 [1.23 , 1.35]	
Subtotal (95% CI)		1878		1892	59.3%	0.62 [0.18 , 2.16]	
Total events:	1174		925				
Heterogeneity: Tau ² = 0	0.78; Chi ² = 5	.70, df = 2	(P = 0.06); I ²	= 65%			
Test for overall effect: 2	Z = 0.75 (P =	0.46)					
1.10.2 Early discharge	> 74 hours						
Waldenström 1987	3	50	5	54	12.2%	0.65 [0.16 , 2.57]	
Boulvain 2004	16	228	21	231	28.5%		
Subtotal (95% CI)		278		285	40.7%	0.75 [0.42 , 1.32]	—
Total events:	19		26				
Heterogeneity: Tau ² = 0	0.00; Chi ² = 0	.05, df = 1	$(P = 0.82); I^2$	= 0%			
Test for overall effect: 2	Z = 0.99 (P =	0.32)					
Total (95% CI)		2156		2177	100.0%	0.80 [0.46 , 1.42]	
Total events:	1193	2150	951	21//	100.0 /0	0.00 [0.40 , 1.42]	
Heterogeneity: $Tau^2 = 0$		44 df = 4		- 58%			$\frac{1}{0.05}$ 0.2 1 5 20
Test for overall effect: 2			(r = 0.03), r	- 5070		Eavou	0.05 0.2 1 5 20 rs early discharge Favours standard
rest for overall effect: A	2 – 0.70 (P –	0.43)				Favou	ravours stalluaru (

Test for subgroup differences: $Chi^2 = 0.07$, df = 1 (P = 0.79), $I^2 = 0\%$

Analysis 1.11. Comparison 1: Early versus standard discharge, Outcome 11: Women breastfeeding (exclusively or partially) at six weeks postpartum

Subtral (95% C1) 1941 316 2.9% 2.49 [1.59, 3.91] Ital evens: 291 19 Heterogeneity: Nat applicable Heterogeneity: Nat applicable Feetorgeneity: Nat applicable Sign (27, 2, 106) Auddenston 1967 (2) 37 49 45 52 9.0% 0.87 [0.72, 1.06] Surp 1900 (3) 63 72 20 25 8.1% 1.09 [0.88, 1.36] Subtral (95% C1) 121 77 17.1% 0.97 [0.78, 1.21] Feat events: 100 65 taterogeneity: Tat ² = 0.01; Chi ² = 2.37, df = 1 (P = 0.12); P = 58% Heterogeneity: Tat ² = 0.01; Chi ² = 2.37, df = 1 (P = 0.12); P = 58% feat ovents: 100 65 1.04 [0.90, 1.12] appon 1997 (5) 43 78 38 97 4.9% 1.41 [1.02, 1.94] Winterbum 2000 (6) 86 121 94 127 1.09% 0.96 [0.82, 1.12] 4.06 [0.99, 1.15] Salvaria 2040 (6) 202 227 1.94 127 1.05 [0.99, 1.15] 1.06 [0.99, 1.15] Salvaria 2040 (6) 100 182 162 1.06 [0.99, 1.15] <th>Study or Subgroup</th> <th>Early disc Events</th> <th>charge Total</th> <th>Standard d Events</th> <th>ischarge Total</th> <th>Weight</th> <th>Risk Ratio M-H, Random, 95% CI</th> <th>Risk Ratio M-H, Random, 95% CI</th>	Study or Subgroup	Early disc Events	charge Total	Standard d Events	ischarge Total	Weight	Risk Ratio M-H, Random, 95% CI	Risk Ratio M-H, Random, 95% CI
Subtoal (95% CI) 1941 316 2.9% 2.49 [1.59, 3.91] trad events: 291 19 19 treerogeneity: Not applicable Fest for overall effect: Z = 3.59 (P < 0.0001)	1.11.1 Trials conducted in	1960s						
Total events: 291 19 Teterogeneity: Not applicable Tess for overall effect: $Z = 3.9$ ($P < 0.0001$) L11.2 Trials conducted in 1905 Waldenström 1987 (2) 37 49 45 52 9.0% 0.87 [0.72, 1.06] Tarty 190 (3) 63 72 20 25 8.1% 1.09 [0.88, 1.36] Total events: 100 65 Teterogeneity: Tart = 0.01; Ch ² = 2.37, df = 1 ($P = 0.12$); $P = 38\%$ Test for overall effect: $Z = 0.25$ ($P = 0.80$) L11.3 Trials conducted in 1995 mith-Hanzhan 1995 (4) 17 35 11 17 2.5% 0.75 [0.46, 1.22] Jagon 1997 (5) 43 78 38 97 4.9% 1.44 [1.02, 1.94] Minterburn 2000 (6) 66 121 94 127 10.9% 0.96 [0.82, 1.12] Jouly 1004 (6) 202 227 194 229 15.2% 1.05 [0.98, 1.13] Jouly 1004 (6) 202 227 194 229 15.2% 1.05 [0.98, 1.13] Jouly 1004 (6) 203 227 194 229 15.2% 1.05 [0.98, 1.13] Jouly 1004 (6) 203 227 194 229 15.2% 1.06 [0.99, 1.15] Total events: 348 337 Teterogeneity: Tart = 0.01; Ch ² = 6.33, df = 3 ($P = 0.10$); $P = 53\%$ Test for overall effect: $Z = 0.53$ ($P = 0.5$) L11.4 Trials conducted in 2005 Sintz Bueno 2005 (6) 190 213 182 217 15.0% 1.06 [0.99, 1.15] Total events: 190 182 L11.5 Trials conducted in 2005 Sintz Bueno 2005 (6) 190 213 182 217 15.0% 1.06 [0.99, 1.15] Total events: 190 182 L11.5 Trials conducted in 2005 Sintz Bueno 2005 (6) 190 213 182 217 15.0% 1.06 [0.99, 1.15] Total events: 190 182 H1.15 Trials conducted in 2015 L11.5 Trials conducted in 2015 L11.5 Trials conducted in 2015 L11.5 Trials conducted in 2015 The for overall effect: $Z = 1.52$ ($P = 0.11$) L11.5 Trials conducted in 2015 The for overall effect: $Z = 0.11$ ($P = 0.01$); $P = 94\%$ Test for overall effect: $Z = 0.01$; ($D^{12} = 17.22$, $df = 1$ ($P < 0.0001$); $P = 94\%$ Test for overall effect: $Z = 0.01$; ($D^{12} = 0.9$ ($P < 0.00001$); $P = 79\%$ Terogeneity: Tart = 0.01; Chi ² = 0.55) Total effect: $Z = 0.04$ ($D^{12} = 0.55$) Terowers standard discharge For ours early discharge For	Hellman 1962 (1)	291	1941	19	316	2.9%	2.49 [1.59 , 3.91]	
Heterogeneity: Not applicable Hese for overall effect: $Z = 3.39$ (P < 0.0001) HIL 2 Trials conducted in 1980s Wademström 1987 (2) 37 49 45 52 9.0% 0.87 [0.72, 1.06] Carty 1990 (3) 63 72 20 25 8.1% 1.09 [0.88, 1.36] Carty 1990 (3) 63 72 20 25 8.1% 1.09 [0.88, 1.36] Subtoal (95% CI) 121 77 17.1% 0.97 [0.78, 1.21] Notal events: 100 65 Heterogeneity: Tau ² = 0.01; Chi ² = 2.37, df = 1 (P = 0.12); P = 58% Test for overall effect: $Z = 0.25$ (P = 0.80) HIL 3 Trials conducted in 1990s Subtoal (95% CI) 43 78 38 97 4.9% 1.41 [1.02, 1.94] Winterbury 2000 (6) 86 121 94 127 10.9% 0.96 [0.82, 1.12] Subtoal (95% CI) 461 470 33.5% 1.04 [0.90, 1.19] Subtoal (95% CI) 461 470 33.5% 1.04 [0.90, 1.19] Subtoal (95% CI) 461 470 33.5% 1.04 [0.90, 1.19] Subtoal (95% CI) 213 217 15.0% 1.06 [0.99, 1.15] Subtoal (95% CI) 213 217 15.0% 1.06 [0.99, 1.15] Subtoal (95% CI) 213 217 15.0% 1.06 [0.99, 1.15] Subtoal (95% CI) 121 27 15.5% 1.07 [1.00, 1.14] Subtoal (95% CI) 121 27 15.5% 1.07 [1.00, 1.14] Subtoal (95% CI) 165 1673 31.5% 0.99 [0.83, 1.18] Heterogeneity: Tau ² = 0.11; Ch ² = 1.22; df = 1 (P < 0.0001); P = 94% Test for overall effect: $Z = 0.51$ (P = 0.30) HIL 3 Trials conducted in 2010S Theorem 10 fere: $Z = 0.21$ (Ch ² = 4.386, df = 9 (P < 0.00001); P = 94% Test for overall effect: $Z = 0.11$ (P = 0.30) Heterogeneity: Tau ² = 0.01; Ch ² = 4.386, df = 9 (P < 0.00001); P = 94% Test for overall effect: $Z = 0.01$ (Ch ² = 4.386, df = 9 (P < 0.00001); P = 94% Test for overall effect: $Z = 0.01$ (Ch ² = 4.386, df = 9 (P < 0.00001); P = 94% Test for overall effect: $Z = 0.01$ (Ch ² = 4.386, df = 9 (P < 0.00001); P = 79% Test for overall effect: $Z = 0.01$ (Ch ² = 4.386, df = 9 (P < 0.00001); P = 79% Favoure standard discharge Favoure strand discharge Favoure standard discharge Favour	Subtotal (95% CI)		1941		316	2.9%	2.49 [1.59 , 3.91]	•
East for overall effect: $Z = 3.99 (P < 0.0001)$ 1.1.2 Trials conducted in 1980s Validenström 1987 (2) 37 49 45 52 9.0% 0.87 [0.72, 1.06] Carty 1990 (3) 63 72 20 25 8.1% 0.99 [0.88, 1.36] Subtola (95% C1) 121 77 17.1% 0.97 [0.78, 1.21] For overall effect: $Z = 0.25 (P = 0.80)$ 1.1.3 Trials conducted in 1990s Smith-Hanaban 1995 (4) 17 35 11 17 2.5% 0.75 [0.46, 1.22] Minterburn 2000 (6) 86 121 94 127 10.9% 0.96 [0.82, 1.12] Subtola (95% C1) 461 70 33.5% 1.04 [0.82, 1.12] Subtola (95% C1) 461 37 38 97 Heterogeneity: Tau ² = 0.01; Ch ² = 6.33, df = 3 (P = 0.10); P = 53% East for overall effect: $Z = 0.53 (P = 0.59)$ 1.1.4 Trials conducted in 2009s Sainz Buene 2005 (6) 190 213 182 217 15.0% 1.06 [0.99, 1.15] Subtola (95% C1) 213 217 15.0% 1.06 [0.99, 1.15] Subtola (95% C1) 10 182 Heterogeneity: Not applicable Fast for overall effect: $Z = 1.52 (P = 0.10)$; P = 94% East for overall effect: $Z = 0.11 (P = 0.0001)$; P = 94% East for overall effect: $Z = 0.11 (P = 0.91)$ 1.1.4 Trials conducted in 2010s Subtola (95% C1) 102 172 Heterogeneity: Tau ² = 0.01; Ch ² = 43.86, df = 9 (P < 0.00001); P = 79% East for overall effect: $Z = 0.11 (P = 0.31)$ 1.16 [0.99, 1.15] Favours standard discharge Favours stan	Total events:	291		19				-
L1.2 Trials conducted in 1980 K Waldenström 1987 (2) 37 49 45 52 9.0% 0.87 [0.72, 1.06] Carty 1990 (3) 63 72 20 25 8.1% 1.09 [0.88, 1.36] Subtotal (95% CI) 121 77 17.1% 0.97 [0.78, 1.21] Oral events: 100 65 Electrogeneity: Tau ² = 0.01; Ch ² = 2.37, df = 1 (P = 0.12); P = 58% 58% Instance of the coveral effect: Z = 0.25 (P = 0.80) 51 17 2.5% 0.75 [0.46, 1.22] Sinith-Hanzhan 1995 (4) 17 35 11 17 2.5% 0.75 [0.46, 1.22] Sinith-Barnaban 1995 (4) 17 35 11 17 2.5% 0.75 [0.46, 1.22] Sinith-Barnaban 1995 (4) 17 35 11 17 2.5% 0.75 [0.46, 1.22] Sinith-Barnaban 1995 (4) 17 35 1.07 [0.96, 1.13] 1.05 [0.98, 1.13] Solutial (95% CI) 461 470 33.5% 1.04 [0.90, 1.15] Solutial (95% CI) 213 217 15.0% 1.06 [0.99, 1.15] Solutial (95% CI) 213 217	Heterogeneity: Not applica	ble						
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Bayoumi 2016 930 1495 1020 1503 16.0% 0.92 [0.87, 0.97] Subtotal (95% CI) 1665 1675 31.5% 0.99 [0.83, 1.18] Fotal events: 1091 1172 Heterogeneity: Tau ² = 0.01; Chi ² = 17.22, df = 1 (P < 0.0001); I ² = 94% Fotal (95% CI) 4401 2755 100.0% 1.04 [0.96, 1.13] Fotal (95% CI) 2020 1775 Heterogeneity: Tau ² = 0.01; Chi ² = 43.86, df = 9 (P < 0.00001); I ² = 79% 0.1 0.2 0.5 1 2 5 10 Fest for overall effect: Z = 0.94 (P = 0.35) Favours standard discharge Favours early discharge Favours early discharge Favours early discharge			170	152	172	15.5%	1.07 [1.00 . 1.14]	L
Subtotal (95% CI) 1665 1675 31.5% 0.99 [0.83, 1.18] For al events: 1091 1172 Heterogeneity: Tau ² = 0.01; Chi ² = 17.22, df = 1 (P < 0.0001); I ² = 94% Test for overall effect: Z = 0.11 (P = 0.91) Fotal (95% CI) 4401 2755 100.0% Fotal (95% CI) 4401 2755 100.0% 1.04 [0.96, 1.13] Fotal events: 2020 1775 100.0% 1.04 [0.96, 1.13] Heterogeneity: Tau ² = 0.01; Chi ² = 43.86, df = 9 (P < 0.00001); I ² = 79% 0.1 0.2 0.5 1 2 5 10 Fest for overall effect: Z = 0.94 (P = 0.35) Favours standard discharge Favours early discharge Favours early discharge	0 ()							ſ
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Heterogeneity: Tau ² = 0.01; Chi ² = 17.22, df = 1 (P < 0.0001); I ² = 94% Fest for overall effect: Z = 0.11 (P = 0.91) Fotal (95% CI) 4401 2755 100.0% 1.04 [0.96, 1.13] Fotal events: 2020 1775 Heterogeneity: Tau ² = 0.01; Chi ² = 43.86, df = 9 (P < 0.00001); I ² = 79% $0.1 0.2 0.5 1 2 5 10$ Fest for overall effect: Z = 0.94 (P = 0.35) Favours standard discharge		1091	2000	1172	10, 5	51.570	0.00 [0.00 ; 1.10]	Ţ
Fest for overall effect: $Z = 0.11 (P = 0.91)$ Fotal (95% CI) 4401 2755 100.0% 1.04 [0.96, 1.13] Fotal (95% CI) 4401 2755 100.0% 1.04 [0.96, 1.13] Fotal events: 2020 1775 Iderogeneity: Tau ² = 0.01; Chi ² = 43.86, df = 9 (P < 0.00001); I ² = 79% Output:			2 df = 10		2 = 94%			
Intercondent conduction Intercondent conductin Intercondent conductin <td>0</td> <td></td> <td></td> <td>1 < 0.0001), 1</td> <td>- 3470</td> <td></td> <td></td> <td></td>	0			1 < 0.0001), 1	- 3470			
Intercondent conduction Intercondent conductin Intercondent conductin <td>Total (95% CI)</td> <td></td> <td>4401</td> <td></td> <td>2755</td> <td>100.0%</td> <td>1.04 [0.96 . 1.13]</td> <td></td>	Total (95% CI)		4401		2755	100.0%	1.04 [0.96 . 1.13]	
Heterogeneity: Tau ² = 0.01; Chi ² = 43.86, df = 9 (P < 0.00001); I ² = 79% $0.1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 +$		2020	1	1775	2733	100.0 /0	1.04 [0.00 ; 1.10]	T
Fest for overall effect: $Z = 0.94$ (P = 0.35)Favours standard dischargeFavours early discharge			6 df - 9 ($I^2 = 70\%$		_H	
Fest for subgroup differences: Chi ² = 15.25, df = 4 (P = 0.004), I ² = 73.8%	0			i < 0.00001);	1 - / 3 /0			
	Test for subgroup difference	es: Chi² = 15	.25, df = 4	4 (P = 0.004),	I ² = 73.8%			

Footnotes

(1) Measured at one month, not reported if exclusive or partial breastfeeding

(2) Measured at two months, not reported if exclusive or partial breastfeeding

(3) Measured at one month, exclusive breastfeeding

(4) Measured at 6 weeks, not reported if exclusive or partial breastfeeding

(5) Measured at one month, 'predominantly' breastfeeding

(6) Measured at one month, any breastfeeding

(7) Measured at 6 weeks, any breastfeeding



Analysis 1.12. Comparison 1: Early versus standard discharge, Outcome 12: Women breastfeeding (exclusively or partially) at six weeks postpartum: mode of birth subgroups

	Early dis	scharge	Standard d	ischarge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI
1.12.1 Vaginal birth							
Hellman 1962 (1)	291	1941	19	316	2.9%	2.49 [1.59 , 3.91]	•
Waldenström 1987 (2)	37	49	45	52	9.0%	0.87 [0.72 , 1.06]	
Carty 1990 (3)	63	72	20	25	8.1%	1.09 [0.88 , 1.36]	
Smith-Hanrahan 1995 (4)	17	35	11	17	2.5%	0.75 [0.46 , 1.22]	←
Gagnon 1997 (5)	43	78	38	97	4.9%	1.41 [1.02 , 1.94]	│ →
Sainz Bueno 2005 (6)	190	213	182	217	15.0%	1.06 [0.99 , 1.15]	
Subtotal (95% CI)		2388		724	42.5%	1.15 [0.90 , 1.47]	
Total events:	641		315				
Heterogeneity: Tau ² = 0.07;	; Chi ² = 36.3	32, df = 5 (P < 0.00001);	$I^2 = 86\%$			
Test for overall effect: Z =	1.12 (P = 0.	26)					
1.12.2 Caesarean birth							
Chiong 2012 (7)	161	170	152	172	15.5%	1.07 [1.00 , 1.14]	
Bayoumi 2016	930	1495	1020	1503	16.0%	0.92 [0.87 , 0.97]	
Subtotal (95% CI)		1665		1675	31.5%	0.99 [0.83 , 1.18]	
Total events:	1091		1172				
Heterogeneity: Tau ² = 0.01;	; Chi ² = 17.2	22, df = 1 (P < 0.0001); I	² = 94%			
Test for overall effect: $Z = 0$	0.11 (P = 0.	91)					
1.12.3 Vaginal or caesarea	n birth or	not specifi	ed				
Winterburn 2000 (6)	86	121	94	127	10.9%	0.96 [0.82 , 1.12]	
3001vain 2004 (6)	202	227	194	229	15.2%	1.05 [0.98 , 1.13]	
Subtotal (95% CI)		348		356	26.1%	1.03 [0.95 , 1.11]	
Total events:	288		288				
Heterogeneity: Tau ² = 0.00;	; Chi ² = 1.2	1, df = 1 (P	= 0.27); I ² =	17%			
Test for overall effect: Z =	0.70 (P = 0.	48)					
Fotal (95% CI)		4401		2755	100.0%	1.04 [0.96 , 1.13]	
Total events:	2020		1775				
Heterogeneity: Tau ² = 0.01;	; Chi ² = 43.	86, df = 9 (P < 0.00001);	I ² = 79%			0.850.9 1 1.1 1.2
est for overall effect: Z =	0.94 (P = 0.	35)				Favours sta	andard discharge Favours early disch
est for subgroup differenc		,	(D = 0.01) 12	00/			5 · · · j · · ·

Footnotes

(1) Measured at one month, not reported if exclusive or partial breastfeeding

(2) Measured at two months, not reported if exclusive or partial breastfeeding

(3) Measured at one month, exclusive breastfeeding

(4) Measured at 6 weeks, not reported if exclusive or partial breastfeeding

(5) Measured at one month, 'predominantly' breastfeeding

(6) Measured at one month, any breastfeeding

(7) Measured at 6 weeks, any breastfeeding

Analysis 1.13. Comparison 1: Early versus standard discharge, Outcome 13: Women breastfeeding (exclusively or partially) at six weeks postpartum: subgroups < 24hr vs > 24 hrs

	Early di	scharge	Standard d	ischarge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI
1.13.1 Early discharge < 2	24 hours						
Sainz Bueno 2005 (1)	190	213	182	217	15.0%	1.06 [0.99 , 1.15]	_
Chiong 2012 (2)	161	170	152	172	15.5%	1.07 [1.00 , 1.14]	
Bayoumi 2016	930	1495	1020	1503	16.0%	0.92 [0.87 , 0.97]	_
Subtotal (95% CI)		1878		1892	46.5%	1.01 [0.90 , 1.15]	
Total events:	1281		1354				Ť
Heterogeneity: $Tau^2 = 0.01$; Chi ² = 22.	80, df = 2 (P < 0.0001); I	² = 91%			
Test for overall effect: Z =	0.21 (P = 0.	84)					
1.13.2 Early discharge > 2	24 hours						
Hellman 1962 (3)	291	1941	19	316	2.9%	2.49 [1.59 , 3.91]	
Waldenström 1987 (4)	37	49	45	52	9.0%	0.87 [0.72 , 1.06]	
Smith-Hanrahan 1995 (5)	17	35	11	17	2.5%		
Boulvain 2004 (1)	202	227	194	229	15.2%	1.05 [0.98 , 1.13]	_
Subtotal (95% CI)		2252		614	29.6%	1.12 [0.77 , 1.63]	
Total events:	547		269				
Heterogeneity: Tau ² = 0.12	; Chi ² = 32.	37, df = 3 (P < 0.00001);	I ² = 91%			
Test for overall effect: Z =	0.59 (P = 0.	56)					
1.13.3 Early discharge rat	nge < 24 ho	urs and >	24 hours				
Carty 1990 (6)	- 63	72	20	25	8.1%	1.09 [0.88 , 1.36]	
Gagnon 1997 (7)	43	78	38	97	4.9%	1.41 [1.02 , 1.94]	
Winterburn 2000 (1)	86	121	94	127	10.9%	0.96 [0.82, 1.12]	
Subtotal (95% CI)		271		249	23.9%	1.10 [0.90 , 1.34]	_
Total events:	192		152				
Heterogeneity: Tau ² = 0.02	; Chi ² = 4.9	7, df = 2 (P	$= 0.08$; $I^2 = 0$	50%			
Test for overall effect: Z =							
		4401		2755	100.0%	1.04 [0.96 , 1.13]	
Total (95% CI)			1775	_,00	10000 /0	10. [0.00, 1.10]	T
Total (95% CI) Total events:	2020						
Total events:	2020 : Chi ² = 43.	86. df = 9 ($I^2 = 79\%$		-	
	; Chi ² = 43.			$I^2 = 79\%$		- Favours stand	0.2 0.5 1 2 5 dard discharge Favours early disc

Footnotes

- (1) Measured at one month, any breastfeeding
- (2) Measured at 6 weeks, any breastfeeding
- (3) Measured at one month, not reported if exclusive or partial breastfeeding
- (4) Measured at two months, not reported if exclusive or partial breastfeeding
- (5) Measured at 6 weeks, not reported if exclusive or partial breastfeeding
- (6) Measured at one month, exclusive breastfeeding
- (7) Measured at one month, 'predominantly' breastfeeding

Analysis 1.14. Comparison 1: Early versus standard discharge, Outcome 14: Women breastfeeding (exclusively or partially) at 12 weeks postpartum

Study or Subgroup	Early dis Events	charge Total	Standard d Events	ischarge Total	Weight	Risk Ratio M-H, Fixed, 95% CI	Risk Ratio M-H, Fixed, 95% CI	
Sainz Bueno 2005	141	213	119	217	100.0%	1.21 [1.03 , 1.41]		
Total (95% CI) Total events:	141	213	119	217	100.0%	1.21 [1.03 , 1.41]	•	
Heterogeneity: Not app Test for overall effect: 2 Test for subgroup differ	Z = 2.39 (P =					Favours st	0.5 0.7 1 1.5 2 andard discharge Favours ear	1 2 rly discharge



Analysis 1.15. Comparison 1: Early versus standard discharge, Outcome 15: Women breastfeeding (partially or exclusively) at six months postpartum

	Early dis	charge	Standard d	ischarge		Risk Ratio	Risk I	Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Rando	m, 95% CI
Waldenström 1987 (1)	0	49	0	59		Not estimable		
Boulvain 2004 (1)	78	220	78	215	48.5%	0.98 [0.76 , 1.26]	-	F
Sainz Bueno 2005 (1)	94	213	76	217	51.5%	1.26 [1.00 , 1.60]	-	-
Total (95% CI)		482		491	100.0%	1.11 [0.87 , 1.43]		
Total events:	172		154					
Heterogeneity: Tau ² = 0.	02; Chi ² = 2	.09, df = 1	(P = 0.15); I ²	= 52%			1 0.2 0.5 1	2 5 10
Test for overall effect: Z	= 0.85 (P =	0.40)				Favours stan	dard discharge	Favours early discharge
Test for subgroup differe	ences: Not a	pplicable						

Footnotes

(1) Any breastfeeding

Analysis 1.16. Comparison 1: Early versus standard discharge, Outcome 16: Infant mortality within 28 days

	Early dis	charge	Standard d	ischarge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Hellman 1962	3	1667	1	217	100.0%	0.39 [0.04 , 3.74]	
Bayoumi 2016	0	1495	0	1503		Not estimable	-
Total (95% CI)		3162		1720	100.0%	0.39 [0.04 , 3.74]	
Total events:	3		1				
Heterogeneity: Not appl	icable						0.01 0.1 1 10 100
Test for overall effect: Z	z = 0.82 (P =	0.41)					Early discharge Standard discharge
Test for subgroup different	ences: Not aj	pplicable					

Analysis 1.17. Comparison 1: Early versus standard discharge, Outcome 17: Infant mortality within one year

	Early dis	charge	Standard d	ischarge		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI	
Hellman 1962	4	1667	1	217	55.1%	0.52 [0.06 , 4.64]		
Waldenström 1987	0	49	1	53	44.9%	0.36 [0.02 , 8.63]		
Total (95% CI)		1716		270	100.0%	0.45 [0.07 , 2.77]		
Total events:	4		2					
Heterogeneity: Chi ² = 0	.04, df = 1 (I	9 = 0.85); I	$^{2} = 0\%$				0.01 0.1 1 10 10	H 00
	est for overall effect: $Z = 0.86$ (P = 0.39) est for subgroup differences: Not applicable						Early discharge Standard discha	arge

Analysis 1.18. Comparison 1: Early versus standard discharge, Outcome 18: Number of contacts with healthcare professionals regarding infant health issues within four weeks of birth

	Early dis	charge	Standard d	ischarge		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95	% CI
Brooten 1994 (1)	25	61	31	61	39.8%	0.81 [0.55 , 1.19]	-	
Gagnon 1997 (2)	12	78	17	97	19.4%	0.88 [0.45 , 1.73]		
Chiong 2012 (3)	30	170	32	172	40.8%	0.95 [0.60 , 1.49]	-	
Total (95% CI)		309		330	100.0%	0.88 [0.67 , 1.16]		
Total events:	67		80				•	
Heterogeneity: Chi ² = 0	.30, df = 2 (I	9 = 0.86); I	$^{2} = 0\%$				0.01 0.1 1	10 100
Test for overall effect: 2	Z = 0.92 (P =	0.36)						andard discharge
Test for subgroup differ	ences: Not a	pplicable						

Footnotes

(1) contacts were acute care visits

(2) contacts were problems pertaining to infant feeding, crying, sleeping, or care of the umbilical cord

(3) measured at 6 weeks, contacts any unscheduled medical consultation

Analysis 1.19. Comparison 1: Early versus standard discharge, Outcome 19: Number of contacts with healthcare professionals regarding maternal health issues within six weeks of birth

	Early dis	charge	Standard d	ischarge		Risk Ratio	Risk Ra	atio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed,	95% CI
Brooten 1994 (1)	6	61	13	61	42.1%	0.46 [0.19 , 1.14]		
Chiong 2012 (2)	16	170	18	172	57.9%	0.90 [0.47 , 1.70]	+	
Total (95% CI)		231		233	100.0%	0.72 [0.43 , 1.20]	•	
Total events:	22		31				•	
Heterogeneity: Chi ² = 1	.40, df = 1 (I	P = 0.24); I	2 = 29%				0.01 0.1 1	10 100
Test for overall effect: 2	Z = 1.28 (P =	0.20)					Early discharge	Standard discharge
Test for subgroup differ	ences: Not a	pplicable						

Footnotes

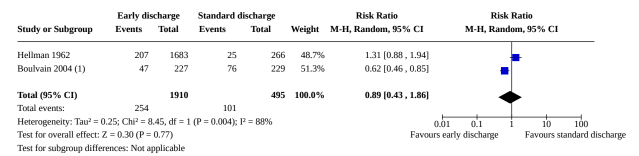
(1) measured at 4 weeks; contacts were acute care visits

(2) measured at 6 weeks, contacts were any unscheduled medical consultation

Analysis 1.20. Comparison 1: Early versus standard discharge, Outcome 20: Women reporting health problems (including perineal pain, perineal infection, breast soreness, breast infection, caesarean wound pain, caesarean wound infection) in the first six weeks postpartum

Study or Subgroup	Early dis Events	scharge Total	Standard d Events	ischarge Total	Weight	Risk Ratio M-H, Fixed, 95% CI	Risk F M-H, Fixed	
Taiba 2012	5	50	60	150	100.0%	0.25 [0.11 , 0.59]		
Total (95% CI)		50		150	100.0%	0.25 [0.11 , 0.59]	•	
Total events:	5		60					
Heterogeneity: Not app	licable					0.0	01 0.1 1	10 100
Test for overall effect:	Z = 3.18 (P =	0.001)				Favours	early discharge	Favours standard discharg
Test for subgroup differ	rences: Not a	pplicable						

Analysis 1.21. Comparison 1: Early versus standard discharge, Outcome 21: Women reporting infant feeding problems



Footnotes

(1) Measured at four weeks

Analysis 1.22. Comparison 1: Early versus standard discharge, Outcome 22: Women satisfied with postnatal care - dichotomous data

	Early dis	scharge	Standard d	ischarge		Risk Ratio	Risk Ratio	Risk of Bias
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI	ABCDEF
Hellman 1962 (1)	1522	1941	272	316	29.7%	0.91 [0.87 , 0.96]		• ? • ? ? •
Waldenström 1987	46	50	22	54	12.5%	2.26 [1.62 , 3.15]		?? 🕂 ?? 🕂
Boulvain 2004	186	217	192	223	28.5%	1.00 [0.92 , 1.07]	_	••••
Sainz Bueno 2005	170	172	113	125	29.3%	1.09 [1.03 , 1.16]	•	? 🖶 🖶 ? ? 🖶
Total (95% CI)		2380		718	100.0%	1.10 [0.95 , 1.29]	•	
Total events:	1924		599				•	
Heterogeneity: Tau ² = 0).02; Chi ² = 4	5.01, df =	3 (P < 0.0000	1); I ² = 93%		⊢ 0.1		⊣ 10
Test for overall effect: 2	Z = 1.27 (P =	0.20)				Favours stand	lard discharge Favours early	discharge
Test for subgroup differ	rences: Not a	pplicable						

Footnotes

(1) Satisfaction measured for in-hospital care only

Risk of bias legend

(A) Random sequence generation (selection bias)

(B) Allocation concealment (selection bias)

(C) Blinding of participants and personnel (performance bias)

(D) Blinding of outcome assessment (detection bias)

(E) Selective reporting (reporting bias)

(F) Other bias

Analysis 1.23. Comparison 1: Early versus standard discharge, Outcome 23: Satisfaction with postnatal care - continuous data

	Earl	y dischar	ge	Standa	ard discha	arge		Std. Mean Difference	Std. Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI	
Carty 1990 (1)	91.55	16.55	49	80.45	20.96	19	20.0%	0.61 [0.07 , 1.15]	-	
Carty 1990 (2)	96.97	11.25	44	80.45	20.96	19	17.8%	1.10 [0.53 , 1.68]	+	
Gagnon 1997	3.6	0.7	78	3	1	97	62.2%	0.68 [0.37 , 0.99]	•	
Total (95% CI)			171			135	100.0%	0.74 [0.50 , 0.98]	•	
Heterogeneity: Chi ² = 1	1.89, df = 2 (P	= 0.39); I	$^{2} = 0\%$							
Test for overall effect:	Z = 6.01 (P <	0.00001)						-10	-5 0 5 10	
Test for subgroup diffe	rences: Not ap	plicable						Favours standa	ard discharge Favours early disch	harge

Footnotes

(1) Intervention arm: early discharge at 25 to 48 hours

(2) Intervention arm: early discharge at 12 to 24 hours

Analysis 1.24. Comparison 1: Early versus standard discharge, Outcome 24: Women who perceive their length of hospital stay as too short

	Early dis	scharge	Standard di	scharge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Yanover 1976	2	41	1	41	100.0%	2.00 [0.19 , 21.21]	
Total (95% CI)		41		41	100.0%	2.00 [0.19 , 21.21]	
Total events:	2		1				
Heterogeneity: Not appl	licable					+ 0.0	
Test for overall effect: Z	z = 0.58 (P =	0.57)				Favours e	early discharge Favours standard discharge
Test for subgroup differ	ences: Not a	pplicable					

Analysis 1.25. Comparison 1: Early versus standard discharge, Outcome 25: Women perceive their length of hospital stay as too long

	Early dis	charge	Standard di	scharge		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Yanover 1976	5	41	9	41	100.0%	0.56 [0.20 , 1.52]	
Total (95% CI)		41		41	100.0%	0.56 [0.20 , 1.52]	
Total events:	5		9				•
Heterogeneity: Not app	licable						0.005 0.1 1 10 200
Test for overall effect: 2	Z = 1.15 (P =	0.25)				Favou	rs early discharge Favours standard dischar
Test for subgroup differ	ences: Not aj	pplicable					

ADDITIONAL TABLES

Table 1. Description of interventions

Study	Early discharge	Standard discharge
Bayoumi 2016	24 hours after delivery (caesarean births only)	72 hours after delivery (caesarean births only)

Boulvain 2004	24 to 48 hours following vaginal births	hospital based postnatal care then dis-
	72 to 84 hours after caesarean births	charge at 4 to 5 days following vaginal births,
	plus home based postnatal care	6 to 7 days following caesareans births
Brooten 1994	'earlier than usual' (mean stay of 3.6 days)	'routine hospital practice' (mean stay of 4.8 days)
	plus minimum of 2 home visits post discharge, plus 10 phone calls to 8 weeks, plus women had phone number to nurse and physician	no routine follow-up care at home post discharge
Burnell 1982	48 hours after delivery (vaginal births only)	conventional hospital stay of 8 to 9
	plus postnatal care at home	days
Carty 1990	Group 1: 12 to 24 hours after delivery (vaginal births only)	4 days after delivery (vaginal births on-
	plus 5 home visits post discharge	ly)
	Group 2: 25 to 48 hours after delivery (vaginal births only)	plus 1 home visit post discharge
	plus 3 home visits post discharge	
Chiong 2012	1 day after delivery (caesarean births only)	2 days after delivery (caesarean births only)
Gagnon 1997	6 to 36 hours after delivery (vaginal births only)	48 to 72 hours after delivery (vaginal births only)
	plus antenatal nursing care at home, at 34 to 38 weeks' gesta- tion, postnatal nursing care by telephone, within 48 hours pp and at 10 days pp, and also at home at 3 and 5 days pp.	plus follow-up as determined by the woman's and infant's physicians.
Hellman 1962	before 72 hours after delivery (vaginal births only)	5 days after delivery (vaginal births only)
Kruse 2021	at 28 hours after delivery (caesarean births only)	48 hours or later after delivery (cae- sarean births only)
McKeever 2002	24 to 36 hours after delivery (vaginal births only)	"standard length of hospitalisa-
	plus up to 3 home visits from community nurses qualified as lactation consultants.	tion" (vaginal births only)
Sainz Bueno 2005	before 24 hours after delivery (vaginal births only)	at least 48 hours after delivery (vagina
	plus home visit by nurse during first 24 to 48 hours	births only)
Smith-Hanrahan 1995	before 60 hours after delivery (vaginal births only)	after 60 hours after delivery (vaginal
	plus telephone call from nurse within 24 hours of discharge leading to a decision to visit or continue to consult by phone; also received phone number for postnatal follow-up service which could be called at any time	births only)
Taiba 2012	72 hours after delivery (caesarean births only)	7 days after delivery (caesarean births only)

Table 1. Description of interventions (Continued)

Thompson 1999	before 36 hours after delivery (vaginal births only)	37-72 hours after delivery (vaginal births only)
Waldenström 1987	24 to 48 hours after delivery (vaginal births only)	6 days after delivery (vaginal births on- ly)
Winterburn 2000	6 to 48 hours after delivery (vaginal and caesarean births) plus community midwife home visits to support breastfeeding	between 48 hours and 7 days after de- livery (vaginal and caesarean births)
Yanover 1976	12 to 48 hours after delivery (vaginal births only)	after 48 hours after delivery (vaginal births only)

pp: postpartum

Table 2. Number of contacts with health professionals regarding infant health issues

Trial	Early discharge	Later discharge	Notes	
Kruse 2021	Median (range): 3 (0-10)	Median (range): 3 (0-9)	Contacts regarding infant and maternal health issues were counted together.	
	54 women	52 women		
Waldenström 1987	452 contacts (49 in-	537 contacts (50 in-	6 month follow-up.	
	fants)	fants)	Number of contacts with child health centre (home visits by CHC nurse, visits to CHC nurse, visits to CHC nurse, visits to CHC paediatrician)	

Table 3. Number of contacts with health professionals regarding maternal health issues

Trial	Early discharge	Later discharge	Notes
Kruse 2021	Median (range): 3 (0-10)	Median (range): 3 (0-9)	Contacts regarding infant and maternal health issues were counted together.
	54 women	52 women	
Boulvain 2004	33/228	48/231	Number of women reporting 1 or more visits to a gynaecologist during the first month.

Table 4. Costs of hospital care in the period immediately following the birth up to the time of discharge

Trial	Early discharge	Later discharge	Difference
Boulvain 2004	CHF 5218	CHF 6772	CHF -1554
	228 women	231 women	
Brooten 1994	USD 7648	USD 10,971	USD -3323
	61 women	61 women	



Table 4. Costs of hospital care in the period immediately following the birth up to the time of discharge (Continued)

Sainz Bueno 2005	USD 382.22	USD 647.67	USD -265.45
	213 women	217 women	

CHF: Swiss francs USD: USA dollars

Table 5. Costs of postnatal care following discharge from hospital in the period up to six weeks after the birth

Trial	Type of cost	Early discharge	Later discharge	Difference
Boulvain 2004	mean cost of all postnatal care	CHF 932	CHF 481	CHF 451
		228 women	231 women	
Boulvain 2004	mean cost of community care (including midwifery, medical and allied health care)	CHF 528 (SD 267)	CHF 234 (SD 273)	CHF 294
		228 women	231 women	
Boulvain 2004	total mean (SD) costs including:	CHF 7798 (SD 6419)	CHF 9019 (SD	CHF -1221
	hospitalisation		4345)	
	 community care (including midwifery, medical and allied health care) 	228 women	231 women	
	 non-medical costs attributable to health care in the first 6 weeks after birth (travel to health care providers, childcare support for siblings) 			
	 costs associated with loss of income if a partner required time off work. 			
Brooten 1994	mean cost of nurse-specialist visits (in hospital and	USD 516	USD 519	USD -3
	at home), home caregiver charges, acute care visits (following discharge) and rehospitalisation charges	61 women	61 women	
Brooten 1994	total mean costs including	USD 8164	USD 11,490	USD -3326
	hospitalisation			
	 nurse-specialist visits (in hospital and at home) 			
	 home caregiver charges 			
	 acute care visits (following discharge) 			
	 rehospitalisation charges 			
Sainz Bueno 2005	combined cost of community care (including ma-	USD 125.24	USD 153.90	USD -28.66
	ternal and neonatal consultations and telephone calls) and maternal and neonatal readmissions	213 women	217 women	

CHF: Swiss francs USD: USA dollars SD: standard deviation

APPENDICES

Appendix 1. Search terms for ICTRP and ClinicalTrials.gov ICTRP



each line was run separately early AND discharge AND birth early AND discharge AND childbirth early AND discharge AND postpartum early AND discharge AND postnatal hospital AND discharge AND caesarean hospital AND discharge AND caesarean timing AND discharge AND caesarean hospital AND discharge AND caesarean **ClinicalTrials.gov** Advanced search

childbirth | Interventional Studies | early discharge

-

childbirth | Interventional Studies | care pathway

postnatal | early discharge

postpartum | early discharge

cesarean | early discharge

WHAT'S NEW

Date	Event	Description
21 May 2021	New search has been performed	Search updated. We included four new trials (Bayoumi 2016; Chiong 2012; Kruse 2021; Taiba 2012), plus three trials that were previously excluded (Burnell 1982; McKeever 2002; Thompson 1999).
21 May 2021	New citation required and conclusions have changed	Early discharge probably leads to a higher risk of infant readmis- sion within 28 days of birth but probably makes little to no differ- ence to the risk of maternal readmission within six weeks post- partum. We are uncertain if early discharge has any effect on the risk of infant or maternal mortality.

HISTORY

Protocol first published: Issue 1, 2001 Review first published: Issue 3, 2002

Date	Event	Description
1 December 2008	New search has been performed	Update. New search identified two additional studies for inclu- sion (Boulvain 2004; Sainz Bueno 2005). Two studies previously awaiting classification were excluded (Burnell 1982a; Thompson 1999a), and a further four studies were also excluded (Escobar



Date	Event	Description
		2001; Lieu 2000; McKeever 2002a; Steel O'Connor 2003). Main re- sults and conclusions remain unchanged.
19 August 2008	Amended	Converted to new review format.

CONTRIBUTIONS OF AUTHORS

For the original review (2002): Stephanie Brown and Rhonda Small undertook the background review of the literature. Stephanie Brown wrote the protocol with input from all review authors. Brenda Argus and Ann Krastev conducted the original literature search to identify trials. Each review author independently evaluated trials for quality and extracted data. Rhonda Small, Brenda Argus and Ann Krastev independently entered the data. Stephanie Brown wrote the text of the review with input from the other review authors.

For the first update (2009): Stephanie Brown and Rhonda Small updated the literature search, independently reviewed new trials, extracted and entered data and updated the text of the review. Brenda Argus, Ann Krastev and Peter Davis reviewed the manuscript.

For this update (2021): Lynn Hampson (Cochrane Pregnancy and Childbirth Information Specialist) updated the literature search, all review authors updated the protocol and agreed the GRADE outcomes for the review update. Ellie Jones and Fiona Stewart independently assessed new trials and previously excluded studies, evaluated trials for risk of bas, extracted and entered data and performed data analyses. Beck Taylor assisted with resolving any disagreement with the data extraction and analysis process. Fiona Stewart and Ellie Jones wrote the text of the review with input from the other review authors Beck Taylor, Stephanie Brown and Peter Davis.

DECLARATIONS OF INTEREST

Stephanie Brown: salary and project support from the Australian National Health and Medical Research Council.

Peter G Davis: salary and project support from the Australian National Health and Medical Research Council.

Beck Taylor: is supported by the UK National Institute for Health Research (NIHR) Applied Research Centre (ARC) West Midlands. The views expressed are those of the author(s) and not necessarily those of the NIHR or the Department of Health and Social Care.

Fiona Stewart: none known

Eleanor Jones: is supported by the National Institute for Health Research (NIHR). The views expressed are those of the author(s) and not necessarily those of the NIHR or the Department of Health and Social Care.

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- La Trobe University, Australia
- Royal Women's Hospital, Melbourne, Australia
- Murdoch Childrens Research Institute, Australia

External sources

• No sources of support provided

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We added in a search of ClinicalTrials.gov and the WHO International Clinical Trials Registry Platform (ICTRP).

We made several changes to the outcomes for the review: maternal and infant mortality at 28 days and one year were added as secondary outcomes; we changed the secondary outcome 'women dissatisfied with their postnatal care' to 'women satisfied with their postnatal care' and the time points for 'Women scoring above the cut-off score indicating probable depression on a well-validated standardized instrument for measuring depression' was changed from six to eight weeks, three months and six months to within six months after the birth.

We also prespecified subgroup analyses for very early discharge (24 hours and under) and early discharge (48 hours and greater) and mode of birth (vaginal birth or caesarean section).

The certainty of evidence has been assessed using the GRADE approach and 'Summary of findings' tables have been added.



INDEX TERMS

Medical Subject Headings (MeSH)

Bias; Breast Feeding [statistics & numerical data]; Depression, Postpartum [epidemiology]; Infant Mortality; *Length of Stay; *Patient Discharge; Patient Readmission [statistics & numerical data]; *Postpartum Period; *Term Birth; Time Factors

MeSH check words

Female; Humans; Infant; Pregnancy