## Children's Mercy Kansas City SHARE @ Children's Mercy

**Clinical Critically Appraised Topics** 

**Critically Appraised Topics** 

4-2020

## Initiating enteral feedings after medical NEC: Summary

Children's Mercy Kansas City

Let us know how access to this publication benefits you

Follow this and additional works at: https://scholarlyexchange.childrensmercy.org/clinical-criticallyappraised-topics

#### **Recommended Citation**

Children's Mercy Kansas City, "Initiating enteral feedings after medical NEC: Summary" (2020). *Clinical Critically Appraised Topics*. 27. https://scholarlyexchange.childrensmercy.org/clinical-critically-appraised-topics/27

This Critically Appraised Topic is brought to you for free and open access by the Critically Appraised Topics at SHARE @ Children's Mercy. It has been accepted for inclusion in Clinical Critically Appraised Topics by an authorized administrator of SHARE @ Children's Mercy. For more information, please contact hlsteel@cmh.edu.

#### Specific Care Question

For the patient in the NICU with medical necrotizing enterocolitis (mNEC) when is the optimal time to restart enteral feedings?

#### **Recommendations from the XXXXX Team**

No recommendation can be made on the timing of the re-initiation of enteral feedings after Stage IIa or IIb NEC. Three cohort studies are included, that show no difference in NEC recurrence or intestinal stricture if feedings are restarted early, and catheter related sepsis is lower when feedings are restarted early. However, the definition of early and late refeeding varied among the studies, which makes the studies inconsistent, and the time frame over which data was collected is wide. Other changes in neonatal care may have occurred to influence the outcomes.

When there is a lack of scientific evidence, standard work should be developed, implemented, and monitored.

#### **Literature Summary**

**Background.** Necrotizing enterocolitis is a condition that affects premature infants. In its most severe form, NEC causes severe inflammation and necrosis of the intestinal mucosa (Kim, 2019, Shenk 2019). It also presents in less severe forms. Medical NEC (mNEC) is when surgery is not required. A staging system, known as Bell staging, has been developed to describe the symptoms:

Name	Bell Stage	Symptoms
Suspected	Stage I	Emesis, abdominal distension, bloody stool
Proven	Stage II a	All the above, plus abdominal tenderness and lack of bowel sounds
Proven	Stage IIb	All the above, plus abdominal cellulitis
Advanced	Stage III	All the above, plus hypotension, pH imbalance, bradycardia,
		neutropenia

*Note*: from (Kim, 2019; Shenk et al., 2019)

Necrotizing enterocolitis is managed by stopping enteral feedings, initiating antibiotics, and continuing other supportive treatment, such as temperature regulation (Hock et al., 2018). There is a lack of standardization of when to re-start enteral feedings after a mNEC event. From a survey sent to 34 Intensive Care Nurseries (n = 22 responses) participating in the Children's Hospitals Neonatal Consortium (O'Donnell et al., 2019):

60% (13/22)- began enteral feeding on the day after antibiotics were completed

23% (5/22)- began enteral feeding on the day of discontinuation of antibiotics

18% (4/22-) starting feeds after resolution of pneumotosis and the return of bowel function

It is unknown if early or late reinitiation in feeding plays a role in NEC recurrence, time to full feeds, growth, or hosptial length of stay. In the calendar year 2017-2018, the NICU at Children's Mercy Kansas City discharged 13 patients with mNEC (Younger, 2019). Variation of timing of the restart of enteral feeds after mNEC is unknown.

The goal of the NICU is to standardize the initiation of enteral feedings after mNEC and to identify process points for feeding infants after mNEC to decrease patient important outcomes such as NEC recurrence, time to full feeds and hospital length of stay. This review will summarize identified literature to answer the specific care question.



**Study characteristics**. The search for suitable studies was completed on December 6, 2019. A. Khmour, MD and Denise Smith, RN, NNP-BC reviewed the 105 titles and/or abstracts found in the search and identified<sup>b</sup> 10 single studies believed to answer the question. A systematic review/meta-analysis was identified by ancestry search and led to two additional? studies that answered the question. After an in-depth review of the fourteen articles<sup>c</sup>, four answered the question (see Figure 1).

**Optimal time to start feedings after mNEC.** Hock et al. (2018) is a systematic review/meta-analysis of two cohort studies that evaluated the timing of re-initiation of enteral feedings after mNEC. The two cohort studies are Bohnhorst et al. (2003) and Brotschi, Baenziger, Frey, Bucher, and Ersch (2009). Hock et al. (2018) reported on the outcomes (a) NEC recurrence, (b) catheter related sepsis, and (c) occurrence of intestinal stricture. Both studies compared shortening of time to initiate enteral feedings after mNEC to a historical cohort. The cohort study Arbra, Oprisan, Wilson, Ryan, and Lesher (2018) is a retrospective cohort that evaluated patients with early or late feeding after mNEC. The data from Arbra et al. (2018) has been added to the meta-analysis in this synthesis (see Table 1 and Figure 2).

Variation in time to feeding. The included studies differed in the definition of early and late enteral feeding.

Study	Arbra et al. (2018)	Bohnhorst et al. (2003)	Brotschi et al. (2009)
Early Feedings	Feeds started < 7 days after diagnosis of NEC, $n = 40$	Feeds started after 3 days without evidence of gas bubbles in the portal vein, n = 26	Feeds started $< 5$ days after NEC diagnosis, $n =$ 30
Dates collected	July 2006 to June 2016	January 1998 to December 2001	January 2000 to December 2006
Delayed Feedings	Feeds started $\geq$ 7 days after diagnosis of NEC, <i>n</i> =98	Per the neonatologist, $n = 18$	Feeds started > 5 days after NEC diagnosis (median 5 days), $n = 17$
Dates collected	July 2006 to June 2016	April 1993 to March 1997	January 2000 to December 2006

#### Summary by Outcome

**NEC recurrence.** Three studies (n = 229) measured NEC recurrence after early initiation of feeds after mNEC (Arbra et al., 2018; Bohnhorst et al., 2003; Brotschi et al., 2009). The studies reported the number of NEC recurrences as counts for early and late reinitiation of enteral feeding, and they are included in the meta-analysis (see Figure 2 and Table 1). The odds of NEC recurrence was not significantly different from restarting feedings later, OR = 0.46, 95% CI [0.15, 1.48].

**Certainty of the evidence for NEC recurrence.** The certainty of the body of evidence was very low based on four factors: *withinstudy risk of bias, consistency among studies, directness of evidence,* and *precision of effect estimates*. The body of evidence was assessed to have very serious risk of bias, very serious imprecision, not serious indirectness and serious inconsistency. Very serious risk of bias was assessed due to design, it is a cohort study, subjects were not randomized into feeding treatments, nor were subjects, providers, or outcome assessors blinded. Imprecision is serious due to the small number of subjects included in the papers. The studies are inconsistent in the definition of both early and late feeding. Furthermore, the date ranges from which the data was pulled was wide. For the Late Feeding group, data was pulled from 1993 to 2006. Time varying confounding occurs when



the intervention received can change over time (Sterne, Higgins, & Reeves, 2014). Research published in the time between 1993 and 2006 may have suggested other changes in the care of the patient with mNEC that may have influenced the outcomes.

**Catheter related sepsis.** Two studies (n = 91) measured catheter related sepsis (Bohnhorst et al., 2003; Brotschi et al., 2009). The studies reported results as counts of catheter related sepsis events, and they are included in the meta-analysis (see Figure 3 and *Table 1*). The odds of catheter related sepsis were not significantly different from restarting feedings later, OR = 0.2, 95% CI [0.01, 3.29].

**Certainty of the evidence for catheter related sepsis.** The certainty of the body of evidence was very low based on four factors: *within-study risk of bias, consistency among studies, directness of evidence,* and *precision of effect estimates*. The body of evidence was assessed to have serious risk of bias, very serious imprecision, not serious indirectness and serious inconsistency. Very serious risk of bias was assessed due to design, it is a cohort study, subjects were not randomized into feeding treatments, nor were subjects, providers, or outcome assessors blinded. Imprecision is serious due to the small number of subjects included in the papers. The studies are inconsistent in the definition of both early and late feeding. Time varying confounding occurs when the intervention received can change over time (Sterne et al., 2014). Research published in the time between 1993 and 2006 may have suggested other changes in the care of the patient with mNEC that may have influenced the outcomes.

**Intestinal stricture.** Three studies (n = 229) measured intestinal stricture (Arbra et al., 2018; Bohnhorst et al., 2003; Brotschi et al., 2009). The studies reported results as counts of stricture occurrence, and they are included in the meta-analysis (see Figure 4 and Table 1). The odds of intestinal stricture were not significantly different from restarting feedings later, OR = 0.15, 95% CI [0.15, 2.37].

**Certainty of the evidence for intestinal stricture.** The certainty of the body of evidence was very low based on four factors: *within-study risk of bias, consistency among studies, directness of evidence*, and *precision of effect estimates*. The body of evidence was assessed to have very serious risk of bias, very serious imprecision, not serious indirectness and serious inconsistency. Very serious risk of bias was assessed because by design, it is a cohort study, subjects were not randomized into feeding treatments, nor were subjects, providers, or outcome assessors blinded. Time varying confounding occurs when the intervention received can change over time (Sterne et al., 2014). Research published in the time between 1993 and 2006 may have suggested other changes in the care of the patient with mNEC that may have influenced the outcomes.

#### Identification of Studies

#### Search Strategy and Results (see Figure 1)

PubMed(1/6/2019)

(mNEC[tiab] OR ((nonoperative\* OR nonsurgical OR medical[tiab]) AND ("Enterocolitis, Necrotizing"[Mesh] OR "Necrotizing Enterocolitis" OR NEC))) and (infants OR infant OR neonate OR "intensive care units, neonatal"[mesh] OR "intensive care, neonatal"[mesh] OR "intensive care nursery") AND ((time OR timing OR early OR duration OR standard OR delay OR restart\* OR resume\* OR reintroduc\* OR initiat\*) AND (enteral OR "enteral nutrition"[mesh] OR feed\* OR refeeding)) Records identified through database searching n = 105Additional records identified through other sources n = 3

Studies Included in this Review



*Studies included from meta-analysis	
Citation	tudy Type
Arbra et al. (2018) Cohort	
Hock et al. (2018) Systematic Revie	ew Meta-Analysis
*Bohnhorst et al. (2003) Cohort	
*Brotschi et al. (2009) Cohort	
Studies Not Included in this Review with Exc	lusion Rationale
Citation	Reason for exclusion
Downard et al. (2012)	Does not answer the question, included in the antibiotic review
Jayanthi, Seymour, Puntis, and Stringer (1998)	Does not answer the question, recommends feeding type for patients with gastroschisis
Kasivajjula and Maheshwari (2014)	Narrative review
Kosloske and Musemeche (1989)	Narrative review
Panigrahi (2006)	Narrative review
Sisk, Lovelady, Dillard, Gruber, and O'Shea (2007)	Does not answer the question, does not address post mNEC
Stringer et al. (1993)	Case series, does not address post mNEC
Thompson and Bizzarro (2008)	Narrative review
Wu, Caplan, and Lin (2012)	Narrative review
Methods Used for Appraisal and Synthesis	
<sup>a</sup> The GRADEpro Guideline Development Tool (GDT)	is the tool used to create the Summary of Findings table(s) for this analysis.
<sup>b</sup> Rayyan is a web-based software used for the ini	itial screening of titles and / or abstracts for this analysis (Ouzzani, Hammady, Fedorowicz
& Elmagarmid, 2017).	
<sup>c</sup> The Preferred Reporting Items for Systematic Re	eviews and Meta-Analyses (PRISMA) flow diagram depicts the process in which literature is

searched, screened, and eligibility criteria is applied (Moher, Liberati, Tetzlaff, & Altman, 2009). <sup>d</sup>Review Manager (Higgins & Green, 2011) is a Cochrane Collaborative computer program used to assess the study characteristics as well as the risk of bias and create the forest plots found in this analysis.

<sup>a</sup>GRADEpro GDT: GRADEpro Guideline Development Tool (2015). McMaster University, (developed by Evidence Prime, Inc.). [Software]. Available from <u>gradepro.org</u>.

<sup>b</sup>Ouzzani, M., Hammady, H., Fedorowicz, Z., & Elmagarmid, A. (2016). Rayyan-a web and mobile app for systematic reviews. *Systematic Reviews*, *5*(1), 210. doi:10.1186/s13643-016-0384-4

<sup>c</sup>Moher D, Liberati A, Tetzlaff J, Altman DG, The PRISMA Group (2009). *P*referred *R*eporting *I*tems for *S*ystematic Reviews and *M*eta-Analyses: The PRISMA Statement. PLoS Med 6(7): e1000097. doi:10.1371/journal.pmed1000097 For more information, visit <u>www.prisma-statement.org</u>.

<sup>d</sup>Higgins, J. P. T., & Green, S. e. (2011). *Cochrane Handbook for Systematic Reviews of Interventions [updated March 2011]* (Version 5.1.0 ed.): The Cochrane Collaboration, 2011.



<b>Ouestion Origina</b>	itor						
Avman Khmou	Avman Khmour, MD						
Denise Smith,	RN, NNP-BC						
Medical Libraria	n Responsible for the Search Strategy						
Keri Swaqqar	t, MLIS, AHIP						
EBP Scholar's Re	sponsible for Analyzing the Literature						
Teresa Bontra	ger, MSN, RN, CPEN						
Kori Hess, Pha	rmD						
Lucy Pappas, I	MS RD CSP LD						
Ashley Wilson,	BSN, RN, CPN						
EBP Team Memb	er Responsible for Reviewing, Synthesizing, and Developing this Document						
Nancy H. Aller	n, MS, MLS, RD, LD, CPHQ						
Acronyms Used in	this Document						
Acronym	Explanation						
CAT	Critically Appraised Topic						
СМН	Children's Mercy Hospital						
EBP	Evidence Based Practice						
mNEC	Medical necrotizing enterocolitis						
NEC	Necrotizing enterocolitis						
NICU	Neonatal intensive care unit						
OR	Odds ratio						
PRISMA	Preferred Reporting Items for Systematic Reviews and Meta-Analyses						
RoB	Risk of bias						
SD	Standard deviation						
sNEC	Surgical necrotizing enterocolitis						
Date Developed	/Updated						
March 2020							





*Figure 1.* Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRIMSA)<sup>c</sup>



Table 1.

## Summary of Findings Table: Re-initiation of Enteral Feeds after mNEC

Certainty assessment						Summary of findings					
							Study event rates (%)			Anticipated absolute effects	
№ of participants (studies) Follow-up	Risk of bias	Inconsistency	Indirectness	Imprecision	Publication bias	Overall certainty of evidence	With Delayed initiation of enteral feeds	With Early re- initiation of enteral feeds after mNEC	Relative effect (95% CI)	Risk with Delayed initiation of enteral feeds	Risk difference with Early re- initiation of enteral feeds after mNEC
NEC recu	rrence	e, lower is b	etter								
229 (3 Cohort)	very serious <sub>a,b</sub>	serious <sup>c,d</sup>	not serious	very serious d	none	⊕⊖⊖⊖ VERY LOW	15/133 (11.3%)	5/96 (5.2%)	<b>OR 0.46</b> (0.15 to 1.48)	113 per 1,000	<b>58 fewer</b> <b>per 1,000</b> (from 94 fewer to 46 more)
Catheter	relate	d sepsis, lo	wer is bett	er							
91 (2 Cohort)	serious <sub>a,b</sub>	serious <sup>c,d</sup>	not serious	very serious d	none		10/35 (28.6%)	5/56 (8.9%)	<b>OR 0.20</b> (0.01 to 3.29)	286 per 1,000	<b>212</b> fewer per <b>1,000</b> (from 282 fewer to 283 more)
Intestina	l Stric	ture								<u> </u>	



Certainty assessment								Sumn	nary of fi	ndings	
229 (3 cohort)	serious <sub>a,b</sub>	serious <sup>c,d</sup>	not serious	very serious	none		6/133 (4.5%)	3/96 (3.1%)	<b>OR 0.59</b> (0.15 to 2.37)	45 per 1,000	<b>18 fewer</b> <b>per 1,000</b> (from 38 fewer to 56 more)

CI: Confidence interval; OR: Odds ratio

#### Notes:

a. It is a cohort study, by design, so it starts at lower level of evidence.

b. Time varying confounding, which is a selection bias occurs due to the wide range of dates in which subjects were enrolled

c. The definition of early and late feeding varied among the studies.

d. Low number of studies that include a low number of subjects.

#### Meta-analyses

	Earl	y	Delay	ed		Odds Ratio		Odds Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M	-H, Random, 95% C	I	
Abra 2018	2	40	12	98	56.3%	0.38 [0.08, 1.77]				
Bohnhorst 2003	2	26	1	18	21.9%	1.42 [0.12, 16.91]				
Brotschi 2009	1	30	2	17	21.8%	0.26 [0.02, 3.09]		• + -		
Total (95% CI)		96		133	100.0%	0.46 [0.15, 1.48]				
Total events	5		15							
Heterogeneity: Tau <sup>2</sup> =	0.00; Ch	i² = 1.0	6, df = 2 (	P = 0.5	9); l² = 09	6			10	100
Test for overall effect:	Z = 1.30	(P = 0.1	9)				0.01 0.1	Early Delayed	10	100

# *Figure 2. Comparison: Early re-initiation of enteral feeds after mNEC versus Delayed initiation of enteral feeds, Outcome: NEC recurrence, lower is better*





*Figure 3. Comparison: Early re-initiation of enteral feeds after mNEC versus Delayed initiation of enteral feeds, Outcome: Catheter related sepsis, lower is better* 



# *Figure 4. Comparison: Early re-initiation of enteral feeds after mNEC versus Delayed initiation of enteral feeds, Outcome: Intestinal Stricture*



#### Arbra et al. (2018)

aracteristics of Study							
Methods	Cohort						
Participants	Participants: NICU patients discharged between 0 and 6 months of age with a discharge diagnosis of NEC over a 10-year study period (July 2006-June 2016) Setting: Tertiary care children's hospital with a 64-bed level IV NICU at the Medical University of South Carolina Number enrolled into study: N = 138 <ul> <li>Group 1, Early feeding (&lt;7 days after NEC diagnosis): n = 40</li> <li>Group 2, Late feeding (≥ 7 days after NEC diagnosis): n = 98</li> </ul> Number completed: N = 138 <ul> <li>Group 1: n = 40</li> <li>Group 2: n = 98</li> </ul> Gender, males: <ul> <li>Group 1: n = 20 (50%)</li> <li>Group 2: n = 49 (50%)</li> </ul> Ethnicity:						
	African American Caucasian Hispanic	<b>Group 1</b> 26 (65%) 12 (30%) 1 (2.5%)	Group 2 54 (55.1%) 37 (37.8%) 6 (6.1%)				
	Other       1 (2.5%)       1 (1.0%)         Age at NEC diagnosis, mean in days:       •         •       Group 1: 19         •       Group 2: 22.6         Inclusion criteria:       •         •       NICU patients discharged between 0 and 6 months of age         •       Discharge diagnosis of NEC over a 10-year study period (July 2006-June 2016)         Exclusion criteria:       •         •       Feeds never restarted after NEC diagnosis         •       Death         Covariates identified:       •         •       Not reported						
Interventions	Both: • Group 1: feed	s restarted <7_days	after NEC diagnosis				



	• Group 2: feeds restarted ≥7 days after NEC diagnosis
Outcomes	<ul> <li>Primary outcome(s):         <ul> <li>NEC recurrence*</li> </ul> </li> <li>Secondary outcome(s)         <ul> <li>Intestinal stricture</li> <li>Mortality</li> </ul> </li> <li>Safety outcome(s): not reported         <ul> <li>*Outcomes of interest to the Children's Mercy Hospital (CMH) Critically Appraised Topic (CAT) Development team</li> </ul> </li> </ul>
Notes	<b>Results:</b> Length of Stay In patients without cardiac disease, length of stay (adjusted for Bell's Stage) was significantly shorter (p<0.01) for Group 1 on linear regression analysis by 30.5 days [95% CI 9.8-51.2] <i>Note</i> : sample size is small.



#### Bohnhorst et al. (2003)

Characteristics of Study	
Methods	Cohort
Participants	<ul> <li>Participants: Infants born at &lt;36 weeks gestational age and admitted to NICU.</li> <li>Setting: Hannover Medical School between January 1, 1998 and December 31, 2001 for group 1; data from retrospective cohort with NEC admitted between April 1, 1993 and March 31, 1997 for group 2.</li> <li>Number enrolled into study: N = 44 <ul> <li>Group 1, Early feeding re-initiation: n = 26</li> <li>Group 2, Historic feeding re-initiation: n = 18</li> </ul> </li> <li>Number completed: N = 44 <ul> <li>Group 2: n = 18</li> </ul> </li> <li>Gender, males: (as defined by researchers)</li> <li>Not reported.</li> </ul> <li>Race / ethnicity or nationality (as defined by researchers): <ul> <li>The study occurred in Germany. The authors did not identify race or ethnicity of the participants.</li> </ul> </li> <li>Age, mean/median in months/years, range/IQR <ul> <li>At least one clinical sign (gastric residuals, abdominal distension, blood in stool) plus gas bubbles in the portal vein or liver parenchyma, pneumatosis intestinalis, and/or free air on ultrasound or radiograph.</li> <li>This definition corresponds to the Bell stage II or higher.</li> </ul> </li> <li>Exclusion criteria: <ul> <li>Not reported</li> </ul> </li> <li>Covariates identified: <ul> <li>Not reported</li> </ul> </li>
Interventions	<ul> <li>Both: Complete cessation of enteral feedings, nasogastric drainage, total parenteral nutrition, and appropriate antibiotic treatment.</li> <li>Group 1: Enteral feedings were reinitiated after 3 consecutive days without evidence of gas bubbles via ultrasound.</li> <li>Group 2: Enteral feedings and advancement performed at the discretion of the attending; usually began at 14 days after onset.</li> </ul>



Outcomes	<ul> <li>Primary outcome(s):         <ul> <li>Early feedings would shorten duration of central venous access</li> </ul> </li> <li>Secondary outcome(s)         <ul> <li>*Enteral feedings restarted at a median of 4 days versus 10 days</li> </ul> </li> <li>Safety outcome(s):         <ul> <li>Not reported</li> </ul> </li> <li>*Outcomes of interest to the CMH CAT development team</li> </ul>
Notes	<ul> <li>Results:</li> <li>Complete enteral feedings were established after 10 days in group 1 compared with 19 days in group 2 (p &lt; .001).</li> <li>Reduction of central line duration (13.5 days vs 26 days; p &lt; .001).</li> <li>In group 1, catheter related septicemia occurred in 18% episodes of NEC compared with 29% in group 2 (P&lt;.01).</li> <li>Time to hospital discharge was 63 vs 69 days (p &lt; .05).</li> </ul>



Brotschi et al. (2009)

Characteristics of Study	
Methods	Retrospective Cohort
Participants	Participants: term and preterm neonates with NEC, Bell stage II Setting: Five tertiary NICUs, over a 7-year, January 2000 to Dec 2006 Number enrolled into study: N = 47 • Group 1: Early feedings, n = 30 • Group 2: Late feedings, n = 17 Number completed: N = 47 • Group 1: n = 30 • Group 2: n = 17 Gender, males: (as defined by researchers) • n = 28 (60%) Race / ethnicity or nationality (as defined by researchers): • The study occurred in Switzerland. The authors did not identify race or ethnicity of the participants. Gestational age, mean in weeks ±SD • Group 1: 32 ±2.8 • Group 2: 31.7 ±3.0 Inclusion criteria: Bell stage II Exclusion criteria: • Bell stage I - definition is vague • Bell stage II- required surgery Covariates identified: • Not reported
Interventions	<ul> <li>Both: The same feeding algorithm was used for all subjects</li> <li>Group 1: Fasted &lt; 5-days to feed re-initiation</li> <li>Group 2: Fasted &gt;5 days to feed re-initiation</li> </ul>
Outcomes	Primary outcome(s):



Notes Results:				
		Fasting period		
	< 5days	>5 days	p value	
NEC recurrence	1	2	.27	
Catheter related sepsis	s 0	5	.004	
Intestinal stricture	1	4	.05	



Hock et al. (2018)

Characteristics of Study	
Design	Quantitative Synthesis (meta-analysis)
Objective	Determine if timing of the initiation of enteral feedings after an episode of Bell stage II NEC influenced the recurrence of NEC
Methods	Protocol and registration.  • Not reported  Eligibility Criteria.  • Human studies  • Assessed the timing of feeds after medical NEC  • Early - starting feeds < 5 days (median)  • Late - starting feeds > 5 days (median)  • Primary outcome was recurrence of NEC  Exclusion criteria  • Feeding protocol was unclear  • Timing of starting feeds after mNEC was unclear • Overlap of data form a study in the same center Information sources.  • MEDLINE 1966 to November 2016 • Google Scholar • Cochrane database Search • Teed OR feeding AND necrotizing enterocolitis Study Selection. • Two authors independently screened, extracted, and analyzed the studies and gave reasons for excluding studies. • All authors achieved consensus when faced with disagreements Data collection process. • All authors achieved consensus when faced with disagreements Data collection process. • All authors achieved consensus when faced with disagreements Data collection process. • All authors achieved consensus when faced with disagreements Data collection process. • The Newcastle-Ottawa scale was employed Summary measures. • Synthesis of results. • RevMan 5.3 to calculate pooled odds ratios with 95% confidence intervals. If the 1 <sup>2</sup> was < 25% a random effects model was used, if ≥ 25% a random effects model was employed.



Results	Study Selection. Number of articles identified: N = 4377 Full-text articles assessed for eligibility: n = 47 o Studies included in qualitative synthesis: n = 2
	<ul> <li>Synthesis of results.</li> <li>There was no significant difference in the recurrence rate of NEC between early and delayed enteral feeding, OR = .61, 95% CI [0.12, 3.16], p = .56</li> <li>There was no significant difference in the occurrence of catheter related sepsis, OR = .2, 95% CI [0.01, 3.29], p = .26</li> <li>There was not significant difference in the occurrence of post NEC strictures, OR = .28, 95% CI [.07, 1.18], p = .08.</li> <li>Risk of bias across studies.</li> </ul>
	Risk of bias was assessed as low on the Newcastle Ottawa scale
Discussion	<ul> <li>Summary of evidence.</li> <li>Starting enteral feeding within 5 days did not increase the incident of recurrent NEC, catheter related sepsis, or occurrence of intestinal stricture.</li> <li>When feedings were started post mNEC, early low volume feedings are suggested</li> <li>It is not known the optimal time to increase to full volume enteral feeds.</li> <li>Limitations.</li> <li>Different definitions for early and late re-initiation of feedings</li> <li>One study included only medical NEC, and the other included mNEC and surgical NEC (sNEC). In the latter study, findings were not reported separately for mNEC and sNEC.</li> </ul>
Funding	Funding.: Conflict of interest was reported as None



References

\*Included in the meta-analysis

- Arbra, C. A., Oprisan, A., Wilson, D. A., Ryan, R. M., & Lesher, A. P. (2018). Time to reintroduction of feeding in infants with nonsurgical necrotizing enterocolitis. *J Pediatr Surg*, 53(6), 1187-1191. doi:10.1016/j.jpedsurg.2018.02.082
- Downard, C. D., Renaud, E., St Peter, S. D., Abdullah, F., Islam, S., Saito, J. M., . . . Aspelund, G. (2012). Treatment of necrotizing enterocolitis: an American Pediatric Surgical Association Outcomes and Clinical Trials Committee systematic review. J Pediatr Surg, 47(11), 2111-2122. doi:10.1016/j.jpedsurg.2012.08.011
- Hock, A. M., Chen, Y., Miyake, H., Koike, Y., Seo, S., & Pierro, A. (2018). Initiation of Enteral Feeding After Necrotizing Enterocolitis. *Eur J Pediatr Surg*, 28(1), 44-50. doi:10.1055/s-0037-1604436
  - \*Bohnhorst, B., Muller, S., Dordelmann, M., Peter, C. S., Petersen, C., & Poets, C. F. (2003). Early feeding after necrotizing enterocolitis in preterm infants. *J Pediatr, 143*(4), 484-487. doi:10.1067/S0022-3476(03)00443-8
  - \*Brotschi, B., Baenziger, O., Frey, B., Bucher, H., & Ersch, J. (2009). Early enteral feeding in conservatively managed stage II necrotizing enterocolitis is associated with a reduced risk of catheter-related sepsis. *J Perinat. Med*, *37*, 701-705. doi:10.1515/JPM.2009.129
- Jayanthi, S., Seymour, P., Puntis, J. W., & Stringer, M. D. (1998). Necrotizing enterocolitis after gastroschisis repair: a preventable complication? *J Pediatr Surg*, 33(5), 705-707. doi:10.1016/s0022-3468(98)90191-9
- Kasivajjula, H., & Maheshwari, A. (2014). Pathophysiology and current management of necrotizing enterocolitis. *Indian J Pediatr, 81*(5), 489-497. doi:10.1007/s12098-014-1388-5
- Kosloske, A. M., & Musemeche, C. A. (1989). Necrotizing enterocolitis of the neonate. *Clin Perinatol*, 16(1), 97-111.
- O'Donnell, B., Brozanski, B. S., Azzuqa, A., Schmidt, H., Maieilo, A., Konnikova, L., . . . Yanowitz, T. D. (2019, November 4-6, 2019). A QI journey to standardize refeeding after treatment for medical necrotizing enterocolitis across a large health system UPMC Division of Newborn Medicine. Paper presented at the CHNC Annual Symposium, Atlanta, GA, USA.
- Panigrahi, P. (2006). Necrotizing enterocolitis: a practical guide to its prevention and management. *Paediatr Drugs, 8*(3), 151-165. doi:10.2165/00148581-200608030-00002
- Sisk, P. M., Lovelady, C. A., Dillard, R. G., Gruber, K. J., & O'Shea, T. M. (2007). Early human milk feeding is associated with a lower risk of necrotizing enterocolitis in very low birth weight infants. *J Perinatol*, *27*(7), 428-433. doi:10.1038/sj.jp.7211758
- Sterne, J. A. C., Higgins, J. P. T., & Reeves, B. C. (2014). A Cochrane Risk of Bias Assessment Tool: for Non-Randomized Studies of Interventions (ACROBAT-NRSI). 1.0.0. Retrieved from http://www.riskofbias.info
- Stringer, M. D., Brereton, R. J., Drake, D. P., Kiely, E. M., Capps, S. N., & Spitz, L. (1993). Recurrent necrotizing enterocolitis. *J Pediatr Surg*, 28(8), 979-981. doi:10.1016/0022-3468(93)90496-8
- Thompson, A. M., & Bizzarro, M. J. (2008). Necrotizing enterocolitis in newborns: pathogenesis, prevention and management. *Drugs, 68*(9), 1227-1238. doi:10.2165/00003495-200868090-00004
- Wu, S. F., Caplan, M., & Lin, H. C. (2012). Necrotizing enterocolitis: old problem with new hope. *Pediatr Neonatol*, 53(3), 158-163. doi:10.1016/j.pedneo.2012.04.001

Younger, D. (2019). ICN Discharge of infants with mNEC 2017-2018. Dataset. Accessed Dec. 6, 2019.

