# Therapeutic benefits of glutamine: An umbrella review of meta-analyses

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Abstract. Glutamine may be an essential amino acid in patients with catabolic disease, as it has been demonstrated that circulating glutamine levels drop during critical illness and following major surgery; this may result in an increase in secondary infection risk, recovery time and mortality rates. However, there is much discrepancy in the literature with regards to randomized controlled studies, and therefore, the present study is an umbrella review of published meta-analyses, conducted to examine the effectiveness of glutamine's role as a therapeutic agent. A search using PubMed, Cochrane Library and CINAHL from January 1st, 1980 to December 31st, 2016 was conducted using the following strategy: 'Glutamine AND (meta-analysis OR systematic review)' and publications were retrieved, which provided quantitative statistical analysis of pooled treatment effects on the relative risks of infectious complications, mortality and length of stay in hospital. A total of 22 meta-analyses were entered into the current umbrella review. As displayed in Tables I, II and III, these analyses are split into three groups, based on different parameters. Of the 19 meta-analyses investigating the effects of infectious complications, 15 identified statistically significant reductions in complications, with relative risks ranging between 0.42 and 0.93. In addition, 12 of the 18 meta-analyses analyzing the length of hospital stays presented statistically significant reductions in the length of stay, with reductions ranging between 0.19 to 4.73 days. Only 4 of the 15 meta-analyses studying mortality effects identified statistically significant reductions in mortality with relative risks ranging between 0.64 and 1.28. Statistically significant heterogeneity was observed in 16 of 22 meta-analyses, and publication bias was observed in five of 11 meta-analyses. Glutamine supplementation for critically ill or surgical patients through parenteral or enteral routes appears to reduce the rate of hospital acquired infectious complications and shortening of the length of stay in hospital. Furthermore, glutamine supplementation appeared to reduce the rate of in-patient mortality, but the majority of meta-analyses did not reach statistical significance. However, researchers must appreciate the positive results with caution in light of the fact that there exists statistically significant heterogeneity for the majority of meta-analyses, and statistically significant publication bias in almost half.

#### Introduction

Glutamine may be a conditionally essential amino acid in patients with catabolic disease as it has been demonstrated that circulating plasma glutamine concentrations drop during critical illness and following major surgery (1). This drop may be due to glutamine's central role in nitrogen transport within the body, with glutamine contributing to both precursors for nucleic acid synthesis, as well as antioxidant defenses through the production of glutathione (2). In addition, glutamine is the preferred fuel for rapidly dividing cells, such an enterocytes in the small intestine and immune cells, such as lymphocytes, monocytes and macrophages. For some patients, the synthesis and release of glutamine from skeletal muscle is insufficient to meet demands, and a deficiency in glutamine may lead to small intestine mucosal injury followed by increased wall permeability and bacterial translocation (3). These negative effects, along with immunosuppression, may all contribute to an increased probability of secondary infection risk, which may impede the recovery time of the patient, or worse, increase in-patient mortality rates.

More than a dozen clinical studies have suggested that the provision of parenteral or enteral glutamine supplementation in both critically ill and surgical patients may improve nitrogen balance, constitutive protein levels and improve immune function, while decreasing infection rates, the length of stay in hospital and mortality rates (4). However, there is much discrepancy regarding comparing single randomized controlled trials on glutamine supplementation's effects on clinical endpoints such as infection rates, length of stay in hospital and mortality (5,6). This discrepancy may be due to differences in study design, selected patient populations, severity of the disease, nutritional status of the patient and/or the differences in glutamine supplementation relative to different forms (free or dipeptide form), doses and methods of administration (parenteral or enteral).

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Given the inconsistency of the existing literature and the insufficient statistical power due to small sample sizes, a pooling of information from individual trials may provide a more precise and accurate estimate of glutamine's role in reducing the rates of infectious complications and mortality. To achieve this result, many investigators have turned to performing a powerful statistical method known as meta-analysis. Meta-analyses are fundamental to provide the highest level of evidence to best inform health care decision-making. Therefore, the purpose and objective of the present paper was to summarize the evidence from previously published meta-analyses regarding the effectiveness of glutamine's role as a therapeutic dietary agent to reduce the occurrence of infectious complications, length of stay in hospital and mortality.

### Materials and methods

An umbrella review was selected for the current study. An umbrella review provides a summary of existing published meta-analyses and systematic reviews and determines whether authors addressing similar review questions independently observe similar results and arrive at similar conclusions (7).

A systematic literature search of PubMed, Cochrane Library and CINAHL from January 1st, 1980 to December 31st, 2016 was conducted using the following search strategy: 'Glutamine AND (meta-analysis OR systematic review)'. Only English language publications were retrieved, which provided quantitative statistical analysis of pooled treatment effects on the relative risks and odds ratios of infectious complications, length of stay in hospital and mortality of critically ill and surgical patients. Meta-analyses or systematic reviews that did not present study specific summary data using a minimum of four randomized controlled trials or which were focused on infants and children were excluded.

For the published papers that were accepted into the review, the following information was extracted and entered into a spreadsheet: Number of publications included in the meta-analysis, number of total subjects, glutamine dose and method of administration, pooled treatment effects on the relative risks and odds ratios of infectious complications, length of stay in hospital and mortality. In addition, papers were assessed for their disclosure of quality assessment, statistical heterogeneity (Cochrane's Q test and I<sup>2</sup> statistic) and publication bias (visual inspection of funnel plots and Egger's or Begg's regression test). As this is a descriptive summary review of meta-analyses, no statistical analyses were performed.

# Results

The initial search strategy identified 106 articles and, following careful review, 26 meta-analyses were retrieved for inclusion into this umbrella review. There were four meta-analyses that were excluded because two were not published in English (8,9), one fell outside the selection criteria, as it was focused on preterm infants (10) and one lacked the necessary methodology and presentation of a well-constructed meta-analysis (11). A flow chart of the meta-analyses selection process is summarized in Fig. 1, and Tables I, II and III provide the detailed analysis of the 22 meta-analyses entered into the umbrella review (12-33).

The results of 19 meta-analyses in Table I indicate that parenteral and/or enteral intakes of 0.3 to 0.45 g/kg/day can provide statistically significantly reductions in the incidence of infectious complications for 15 of the 19 meta-analyses with relative risks of all 19 ranging between 0.42 and 0.93. Statistically significant heterogeneity was only observed in two of the 19 meta-analyses, however publication bias was observed in four of the nine meta-analyses that chose to use a funnel plot analysis and/or Egger or Begg's regression tests. Fig. 2 displays the bar graph that represents the frequency of the meta-analyses relative rates for the incidence of infectious complications.

The results of 18 meta-analyses in Table II demonstrate that parenteral and/or enteral intakes of 0.3 to 0.45 g/kg/day can provide statistically significantly reductions in the length of stay in hospital for 12 of the 18 meta-analyses. The range for the reduction in length of stay in hospital for all 12 meta-analyses was between 0.19 to 4.73 days. However, statistically significant heterogeneity was only observed in 16 of the 18 meta-analyses. Publication bias was observed in only one of the nine meta-analyses, which chose to use either funnel plot analysis and/or Egger or Begg's regression test. Fig. 3 displays the bar graph that represents the frequency of the meta-analyses average reduction in length of stay in hospital.

The results of 15 meta-analyses in Table III indicate that parenteral and/or enteral intakes of 0.3 to 0.42 g/kg/day provided statistically significant reductions in mortality for only four of the 15 meta-analyses with relative risk for all 15 meta-analyses ranging between 0.64 to 1.28. Statistically significant heterogeneity was not observed in any of the 15 meta-analyses, and publication bias was observed in only one of the 10 meta-analyses, which chose to use funnel plot analysis and/or Egger or Begg's regression test. Fig. 4 displays the bar graph that represents the frequency of the meta-analyses relative rates for mortality.

## Discussion

With regards to the incidence of infectious complications among critically ill and surgical patients, 15 of the 19 meta-analyses (79%) presented statistically significant reductions. The rate of infection for all 19 meta-analyses demonstrated a reduction ranging from 7 to 58%. Just over half of the meta-analyses (10 of 19) identified reductions of between 10 and 30%, and one-third of the meta-analyses (seven of 19) found reduction of between 30 and 50%. It should be noted that four of the nine meta-analyses presented with significant publication bias, but more troubling is the fact that only nine of the 19 meta-analyses (47%) assessed for publication bias. Publication bias occurs because small studies with null results tend not to be published, and this problem is referred to as the 'file drawer problem'. As published studies are more likely to report positive research outcomes than unpublished ones, the significance of the effect size of the weighted average of the published studies is overestimated, and this can potentially bias the results of the meta-analysis. On a positive note, only one of the 15 meta-analyses that identified statistically significant reductions in infectious complications presented with statistically significant heterogeneity.



Table I. Summary of the meta-analyses on infectious complications that qualified for the present umbrella review on glutamine.

Meta-analysis authors and date	Patient type	Number of studies in meta- analysis	Number of subjects in meta analysis	Average glutamine dose and admin- istration	Main findings of meta- analysis RR and P-value	Q-test P-value	I <sup>2</sup> statistic %	Funnel plot or Egger or Begg's test P-value	(Refs.)
Novak et al 2002	Mixed	7	326	0.30 PN and EN	0.80 P=0.03	0.43	NR		(12)
Avenell 2006	Mixed	19	1,370	NR	0.76 P=0.002	0.26	16	0.03	(13)
Avenell 2009	Mixed	24	1,776	0.42 PN and EN	0.81 P=0.003	0.06	33	+ve funnel plot	(14)
Bollhalder et al 2013	Mixed	30	2,014	0.42 PN and EN	0.83 P=0.009	0.22	17	0.36	(15)
Chen et al 2014	Mixed	15	2,862	0.40 PN and EN	0.85 P=0.02	0.01	51	NS	(16)
Crowther et al 2009	Critically ill	7	274	0.40 PN and EN	0.77 P=0.03	0.46	0		(17)
Asrani et al 2013	Critically ill	8	365	0.36 PN and EN	0.58 P=0.009	0.29	18		(18)
Zhong et al 2013	Critically ill	4	186	0.40 PN and EN	0.59 P=0.006	0.37	4		(19)
Tao <i>et al</i> 2014	Critically ill	33	2,303	0.38 PN and EN	0.79 P<0.000	NR	8	+ve funnel plot	(20)
Wischmeyer et al 2014	Critically ill	12	1,264	0.35 PN and EN	0.86 P=0.09	0.06	43	0.05	(21)
Oldani et al 2015	Critically ill	15	2,795	0.38 PN and EN	0.88 P=0.11	0.004	56	NS	(22)
Yong et al 2016	Critically ill	6	243	NR PN and EN	0.62 P=0.002	0.45	0	0.093	(23)
van Zanten et al 2015	Critically ill	5	776	0.38 EN only	0.93 P=0.39	0.40	0	0.23	(24)
Jiang et al 2004	Surgical	10	355	0.45 PN only	0.42 P=0.002	0.97	NR		(25)
Zheng et al 2006	Surgical	5	215	0.32 PN only	0.76 P=0.04	0.94	NR		(26)
Wang et al 2010	Surgical	10	447	0.34 PN only	0.59 P=0.003	0.65	0		(27)
Yue et al 2013	Surgical	13	653	0.31 PN only	0.52 P=0.002	0.83	0		(28)
Kang et al 2015	Surgical	8	872	0.34 PN only	0.67 P=0.008	0.06	49		(29)
Sandini <i>et al</i> 2015	Surgical	13	1,011	0.37 PN and EN	0.64 P=0.087	0.14	32		(30)

EN, enteral; NR, not reported; NS, not significant; PN, parenteral; RR, relative risk.

Meta-analysis authors and date	Patient type	Number of studies in meta- analysis	Number of subjects in meta analysis	Average glutamine dose and admin- istration	Reduction of hospital length of stay in days	Q-test P-value	I <sup>2</sup> statistic %	Funnel plot or Egger or Begg's test P-value	(Refs.)
Novak <i>et al</i> 2002	Mixed	10	541	0.30 PN and EN	2.6 P<0.05	0.002	NR		(12)
Bollhalder <i>et al</i> 2013	Mixed	30	2,109	0.42 PN and EN	2.35 P=0.001	0.001	86	0.54	(15)
Chen <i>et al</i> 2014	Mixed	14	2,777	0.40 PN and EN	1.48 P=0.24	<0.0001	96	NS	(16)
Crowther et al 2009	Critically ill	12	552	0.40 PN and EN	0.42 NS	NR	36		(17)
Asrani <i>et al</i> 2013	Critically ill	9	390	0.36 PN and EN	1.35 P=0.17	0.0001	78		(18)
Lin et al 2013	Critically ill	4	155	0.44 EN only	3.37 NS	0.02	NR		(31)
Tao <i>et al</i> 2014	Critically ill	36	2,963	0.38 PN and EN	3.46 P<0.0001	NR	63	NS	(20)
Wischmeyer <i>et al</i> 2014	Critically ill	11	639	0.35 PN and EN	2.56 P=0.02	0.002	63	0.87	(21)
Oldani et al 2015	Critically ill	19	2,635	0.38 PN and EN	1.73 P=0.09	0.02	44	NS	(22)
Yong <i>et al</i> 2016	Critically ill	7	303	NR PN and EN	3.89 P<0.0001	0.38	6	0.76	(23)
van Zanten <i>et al</i> 2015	Critically ill	7	583	0.38 EN only	4.73 P=0.02	0.05	52	0.48	(24)
Mottaghi <i>et al</i> 2016	Criticallyl	6	1,286	0.42 EN only	0.19 NS	<0.01	95	+ve funnel plot	(32)
Jiang et al 2004	Surgical	8	275	0.45 PN only	3.25 P=0.0001	0.0003	NR		(25)
Zheng et al 2006	Surgical	6	291	0.32 PN only	3.55 P<0.0001	<0.0001	NR		(26)
Wang et al 2010	Surgical	11	377	0.34 PN only	3.84 P<0.0001	<0.0001	84		(27)
Yue et al 2013	Surgical	12	525	0.31 PN only	3.33 P<0.0001	<0.0001	88		(28)
Kang et al 2015	Surgical	4	729	0.34 PN only	1.72 P=0.03	<0.0001	91		(29)
Sandini et al 2015	Surgical	13	1,000	0.37 PN and EN	2.67 P<0.0001	0.0001	86	NS	(30)

Table II. Summary of the meta-analyses on length of stay in hospital that qualified for the umbrella review on glutamine.

In regard to the length of stay in hospital, 12 of the 18 meta-analyses (67%) presented statistically significant decreases in which the time in hospital decreased between

0.19 to 4.73 days. There were  $\sim$ two-thirds of the meta-analyses (11 of 18) that demonstrated a 2 to 4 day reduction in the length of stay in hospital. However, it should be noted that 16 of the



Table III. Summar	y of the meta-anal	yses on mortalit	y that c	ualified for this	umbrella r	eview on glutamine.
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Meta-analysis authors and date	Patient type	Number of studies in meta- analysis	Number of subjects in meta analysis	Average glutamine dose and admin- istration	Main findings of meta- analysis RR and P-value	Q-test P-value	I <sup>2</sup> statistic %	Funnel plot or Egger or Begg's test P-value	(Refs.)
Novak et al 2002	Mixed	14	751	0.30 PN and EN	0.78 NS	0.99	NR		(12)
Avenell 2006	Mixed	15	1,449	NR	0.81 P=0.07	0.33	11	NS	(13)
Avenell 2009	Mixed	21	1,964	0.42 PN and EN	0.84 P=0.17	0.16	23	NS	(14)
Bollhalder <i>et al</i> 2013	Mixed	22	2,126	0.42 PN and EN	0.89 P=0.15	0.53	0	0.61	(15)
Chen et al 2014	Mixed	17	3,383	0.40 PN and EN	1.01 P=0.87	0.26	17	NS	(16)
Heyland et al 2003	Critically ill	4	397	NR	0.71 P=0.04	0.91	NR		(33)
Crowther <i>et al</i> 2009	Critically ill	8	572	0.40 PN and EN	0.88 P=0.49	0.45	0		(17)
Asrani et al 2013	Critically ill	9	380	0.36 PN and EN	0.70 P=0.001	0.99	0		(18)
Zhong et al 2013	Critically ill	4	190	0.40 PN and EN	0.74 P=0.01	0.84	0		(19)
Tao <i>et al</i> 2014	Critically ill	36	3,454	0.38 PN and EN	0.89 P=0.10	NR	22	+ve funnel plot	(20)
Wischmeyer <i>et al</i> 2014	Critically ill	24	2,312	0.35 PN and EN	0.88 P=0.10	0.58	0	0.57	(21)
Oldani et al 2015	Critically ill	24	2,834	0.38 PN and EN	0.93 P=0.33	0.31	11	NS	(22)
Yong et al 2016	Critically ill	6	243	NR PN and EN	0.64 P=0.02	0.93	0	0.09	(23)
van Zanten <i>et al</i> 2015	Critically ill	10	1,022	0.38 EN only	0.94 P=0.74	0.26	21	0.18	(24)
Mottaghi <i>et al</i> 2016	Critically ill	8	1,327	0.42 EN only	1.28 P=0.07	0.62	0	NS	(32)

EN, enteral; NR, not reported; NS, not significant; PN, parenteral; RR, relative risk.

18 meta-analyses (89%) presented with statistically significant heterogeneity and this weakens the clinical certainty of this effect. Ideally the studies combined into any given meta-analysis should all have used the same experimental protocols; however increased heterogeneity is inevitable due to the wide variation in study design. Differences in study design include number of subjects, age, sex, body mass index, nutritional status (plasma glutamine concentration at admission), total energy intake, type of disease or surgery, disease severity, dosage, timing, duration, chemical form (L-glutamine, alanyl-L-glutamine or glycyl-L-glutamine) and route of glutamine administration (parenteral vs. enteral). On a positive note, only one of the nine meta-analyses presented with statistically significant publication bias, but again this is problematic as only nine of the 18 meta-analyses (50%) assessed for publication bias.

In regard to mortality rate, 10 of the 15 meta-analyses (67%) identified a reduction that ranged between 10 and 30%, however, only four of the 15 meta-analyses (27%) demonstrated statistically significant reductions. More troubling is



Figure 1. Flow chart of meta-analysis selection.



Figure 2. Frequency of the meta-analyses relative rates for the incidence of infectious complications.

the fact that these four meta-analyses represented four of the six that possessed <10 clinical studies in their make-up (4, 4, 6 and 9 respectively). Meanwhile, nine of the 11 meta-analyses with non-significant findings were made up of between 10 to 36 clinical studies, and so there appears to be a bias towards meta-analyses presenting statistical significance when using <10 clinical trials. This relationship was not observed in the infectious complications or length of stay in hospital umbrella reviews. However, heterogeneity and publication bias was never found to be an issue, as none of the meta-analyses presented statistically significant heterogeneity, and only one of the 10 meta-analyses suggested a potential for publication bias.

The apparent effect of glutamine supplementation to reduce the incidence of infectious complications in critically ill and



Figure 3. Frequency of the meta-analyses average reduction in length of stay in hospital.



Figure 4. Frequency of the meta-analyses relative rates for mortality.

surgical patients is probably due to the fact that glutamine is the preferred fuel for enterocytes for maintaining gastrointestinal structure and decreasing gut mucosal atrophy, which could otherwise lead to increased intestinal permeability and bacterial translocation (3). In addition, glutamine positively affects lymphocyte proliferation, cytokine production, macrophage phagocytosis and the secretion of intestinal immune globulin IgA, which further reduces bacterial adhesion and prevents translocation (34). Therefore, glutamine serves a critical role in the protection of the intestinal immune barrier and leads to increased resistance to bacterial translocation and, hence, reduces infectious complication rates. This fact is supported by a meta-analysis that presented beneficial changes in the markers of intestinal inflammation and mucosal permeability with abdominal surgery patients (35). Finally, glutamine supplementation enhances cellular heat shock protein production (HSP70 and HSP72) and glutathione expression, which both protect cells and enhance cell survival. Heat shock proteins are involved in the repair and removal of damaged proteins, and glutathione reacts directly with reactive oxygen species in order to prevent oxidative damage (36). Collectively, both mechanisms of action reduce the likelihood of cellular apoptosis.

If glutamine supplementation can reduce the incidence of hospital acquired infectious complications, then, when averaged over a large number of patients, this can result in a shorted length of stay in hospital. If a critically ill or surgical patient does acquire an infectious complication, this will prolong that patient's length of stay in hospital and require



more resources. Hospital acquired infections are expensive to treat, and it is estimated that hospital acquired infections can prolong a patient's length of stay by at least 4 days; this can have a significant financial impact on hospitalization cost (27). It has been observed that the additional cost of supplemented glutamine treatment is more than completely offset by cost savings in hospital care (37).

Low plasma glutamine concentrations in patients admitted to hospitals are an independent predictor of premature death, but it is unclear whether the decline of circulating glutamine contributes to death, or whether it is a simple marker of disease severity (38). Assuming that supplemental glutamine's primary mechanism of action is in the prevention of hospital acquired infections, then it is understandable why there were no significant changes observed in mortality rates. To begin with, the mortality rate for patients admitted to hospital as inpatients is ~2% (39). Furthermore, hospital acquired infections occur in ~4% of all admitted patients and only ~6% of those patients with hospital acquired infections die during their hospitalization (40,41). From these data, it can be estimated that the mortality rate due to hospital acquired infections for all patients admitted to hospitals is ~0.25%, which represents only an eighth of all in-patient mortality rates. Therefore, for the majority of meta-analyses that did not identify a significant difference in mortality rates, this is probably due to the fact that the majority of clinical studies used in these meta-analyses were too small and too underpowered to detect these small positive changes in mortality rates.

Although enteral glutamine supplementation should benefit intestinal enterocyte function and overall gut barrier structure better than parenteral glutamine supplementation, it has been suggested that there is a greater treatment effect observed for parenteral compared with enteral glutamine supplementation (12,14,16,18). This is thought to be due to the higher bioavailability of glutamine through parenteral administration, whereas enteral glutamine is used by the gut, therefore the immune system outside the gut would not have been affected by enteral glutamine supplementation. However, only one of the eight meta-analyses that performed a subgroup analysis on administration type presented a significant difference between parenteral and enteral administration, and this meta-analysis found that the length of stay in hospital was significantly shortened for the enteral route (4.5 days less for enteral vs. 2.5 days less for parenteral) (23). To confuse matters more, the only two meta-analyses on clinical studies that used enteral glutamine supplementation with critically ill patients demonstrated completely opposite effects on length of stay in hospital with one presenting a 4.73 day reduction and the other with a 0.19 day reduction (it should be noted that these two findings represent the two extremes from the 18 meta-analyses in this category) (24,32). Although both of these meta-analyses examined clinical studies using enteral feeding in critically ill patients only, there was only one common clinical study used by both meta-analyses of the combined total of 12 different clinical studies. The fact that there appears no real difference in outcomes between enteral and parenteral should make sense, considering that many of the potential protective mechanisms of glutamine supplementation given either enterally or parenterally overlap and are quite similar.

Four of six meta-analyses that performed subgroup analysis on high vs. low dose glutamine supplementation observed that the mortality rate was lower with high dose compared to low dose glutamine supplementation, but the differences between groups were never statistically significant (12,14-16,22,30). Of the three meta-analyses that performed subgroup analyses comparing critically ill vs. surgical patients, it also appears that there was no significant difference between these two groups in their response to glutamine supplementation (12,15,16).

In regard to side effects, only 11 of the 22 meta-analyses (50%) addressed this topic with six of them noting that no side effects were observed (12-14,19,24,26), and the remaining five meta-analyses all mentioned the REDOX study that indicated that critically ill patients with glutamine doses of >0.5 g/kg/day had higher urea concentrations and higher mortality rates (20-22,32). However, it has been noted that there were problems with the REDOX study as 30% of the patients in the glutamine supplementation group were shown to have baseline renal failure at admission, which is a very common exclusion criteria for the majority of glutamine supplementation clinical trials (16,21). In the REDOX post-hoc analysis the authors of the study concluded that high-dose glutamine supplementation was only associated with increased mortality in critically ill patients with multi organ failure (42).

Using mostly Jadad or Cochrane quality assessment scales, 18 of the 22 meta-analyses obtained quality assessment scores and the average percentages of high and low quality clinical studies used in any given meta-analysis was 30.4 and 13.8%, respectively. Although none of the meta-analyses specified that clinical studies were excluded from their analysis, more recently published meta-analyses may have excluded low quality studies from the outset without specifying that they were excluded. For the majority of the meta-analyses (11 of the 22), the number of clinical studies entered for analysis ranged between 10 and 20. There were five meta-analyses that used <6 clinical studies, and one meta-analysis published in 2014 used 53 clinical studies, and these six analyses represent 43% of the 125 unique clinical studies used by all 22 meta-analyses (20). Of the 125 unique clinical studies used by the 22 meta-analyses in this umbrella review, 57 (46%) were used only once. This potentially suggests that there are many clinical studies of such poor quality that they may have made the inclusion criteria by one meta-analysis but excluded by all the others. Or it may simply be due to the fact that a particular meta-analysis may have had a very specific focus [acute pancreatitis (18,19), burn patients (31) or gastrointestinal tumor patients (29)] and therefore the clinical studies used by this meta-analysis would have been specifically selected by this one meta-analysis and no other.

This umbrella review has certain limitations. One limitation is that only three indexing systems were searched, and thus it is possible that some meta-analyses were not identified. Another limitation is that only one author performed the search and selection of the meta-analyses included in this umbrella review. And finally, as in all literature reviews, the quality of this umbrella review is directly related to the quality of the included meta-analyses, which are dependent upon the quality of the individual clinical studies used to conduct the meta-analysis.

In conclusion, this umbrella review supports the statement by the American Society for Parenteral and Enteral Nutrition that suggests that parenteral glutamine supplementation should probably be given early and in doses between 0.2 and 0.5 g/kg/day (43). This statement is supported by the fact that a majority of the meta-analyses reviewed identified significant positive benefits for glutamine supplementation to reduce the rate of hospital acquired infectious complications, and shortening of the length of stay in hospital. In addition, glutamine supplementation appeared to reduce the rate of hospital mortality, but for the majority of these meta-analyses they did not reach statistical significance. However, it must be appreciated that all these positive results with caution in light of the fact that there exists statistically significant heterogeneity in the length of stay in hospital meta-analyses, and statistically significant publication bias in the infectious complications meta-analyses. Also, the lack of quality assessment of published clinical studies is problematic, as clinical studies of very low quality may have been included in these meta-analyses, which can therefore potentially bias their overall outcomes. Despite the problems of heterogeneity, publication bias and quality assessment, the potential benefits observed in this umbrella review strongly suggests that parenteral and enteral glutamine supplementation as a component of nutritional support can be considered as an approach to improve the outcomes of critically ill and surgical patients. Finally, in regards to determining if there is beneficial effect on reducing hospital mortality rates with glutamine supplementation, more well-designed multicenter randomized controlled trials using large populations are required.

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