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Review

Haloperidol as prophylactic treatment for postoperative nausea and vomiting: Systematic literature review[☆]

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ABSTRACT

Introduction: The effectiveness of haloperidol for the prophylaxis of postoperative nausea and vomiting (PONV) has been proven in prior trials summarized by Buttner in 2004. New evidence has surfaced since then. Our objective is thus to update the current knowledge on the topic. A systematic review and a meta-analysis were performed, in order to determine the effectiveness and safety of the use of haloperidol as prophylaxis for PONV.

Methodology: The systematic search, the selection of relevant articles, the extraction of data, the critical analysis of the primary studies, the comparisons and analyses were all based on the recommendations of the Cochrane Collaboration and using RevMan5 software.

Results: Ten controlled clinical trials published between 1962 and 2010, that included 2,711 patients, met the selection criteria. As compared against droperidol (RR: 0.97; 95% CI: 0.52–1.79) and against ondansetron (RR: 1.24; 95% CI: 0.66–2.35), no differences were found in terms of effectiveness after 24 h. A protective effect against PONV associated with the use of haloperidol at varying doses, routes of administration and timing of administration was observed as compared with placebo. No significant increases in adverse events have been reported.

Discussion: This systematic review supports the effectiveness of haloperidol as prophylactic treatment of PONV. No statistically significant differences were found as compared against ondansetron or droperidol.

Conclusions: Haloperidol is an effective prophylactic drug for PONV.

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Haloperidol como profilaxis para náuseas y vómito postoperatorios: revisión sistemática de la literatura

RESUMEN

Palabras clave: Haloperidol

Vómitos Náusea Ensayo clínico Introducción: La efectividad del haloperidol en la profilaxis para náuseas y vómito postoperatorios (NVPO) ha sido demostrada en estudios previos resumidos en 2004 por Buttner. Desde entonces ha surgido nueva evidencia, por lo cual nuestro objetivo es actualizar el estado presente del conocimiento en este tema. Se realizó una revisión sistemática y metaanálisis con el fin de aproximarnos a la efectividad y a la seguridad del uso de haloperidol en la profilaxis de NVPO.

Metodología: La búsqueda sistemática, la selección de artículos relevantes, la extracción de datos, el análisis crítico de los estudios primarios, las comparaciones y los análisis se realizaron con base en las recomendaciones de Cochrane Collaboration y a través del software RevMan5.

Resultados: Diez experimentos clínicos controlados, publicados entre 1962 y 2010, que incluyen 2.711 pacientes, cumplen los criterios de selección. Comparado con el droperidol (RR: 0,97; IC 95%: 0,52-1,79) y con el ondansetrón (RR: 1,24; IC 95%: 0,66-2,35), no se encontraron diferencias en la efectividad a las 24 h. Se evidencia un efecto protector contra NVPO asociado al uso de haloperidol en diferentes dosis, vías de administración y momentos de administración al comparar frente a placebo. No hay reporte de aumento de efectos adversos de forma significativa.

Discusión: La efectividad de haloperidol como profilaxis de NVPO queda sustentada por esta revisión sistemática sin que se logren identificar diferencias estadísticamente significativas cuando se compara con el ondansetrón o el droperidol.

Conclusiones: El haloperidol es un medicamento efectivo y seguro para la profilaxis de NVPO. © 2012 Sociedad Colombiana de Anestesiología y Reanimación. Publicado por Elsevier España, S.L. Todos los derechos reservados.

Introduction

Postoperative nausea and vomiting (PONV) are a frequent problem associated with the administration of anesthesia and sedation resulting in patient dissatisfaction, delayed discharge and unplanned admissions. Other complications described include surgical wound dehiscence and hematoma, hydro-electrolytic imbalance, bronchoaspiration of gastric contents and esophageal rupture. ^{2,3}

Some of the risk factors independently related to PONV are: gender, non-use of tobacco, a history of PONV and the administration of opiates during the perioperative period. The frequency of these complications varies, depending on the type and duration of surgery, the type of anesthesia, anesthetic drugs and management of postoperative pain. 6

Haloperidol is an antagonist of the D2 dopaminergic receptors used in psychiatry and in the medical – surgical management of delirium for over 40 years. It belongs to the group of butyrophenones which are potent antiemetic as is droperidol, one of the most commonly used and cost-effective medicines for the management of PONV, prior to the warning issued by the Food and Drug Administration in 2003, regarding its relationship to the development of cardiac arrhythmia. Here is now a renewed interest in the use of haloperidol for the management of PONV, because it is an effective, safe and low cost alternative, with theoretical advantages such as an extended half-life with a delayed potential protective effect. 11,12

A meta-analysis published by Buttner et al. in 2004⁷ evaluated the effectiveness of haloperidol under varying scenarios, including PONV, showing adequate effectiveness with no relationship between the dose used and the scope of the effect. Since then, new evidence has been found^{13,14} evaluating haloperidol's effectiveness^{15–21} and so we considered it appropriate to undertake a new systematic review to assess the effectiveness and safety of haloperidol as a prophylactic agent in PONV.

Objectives

Main objective

 To estimate the effectiveness of haloperidol for the prophylaxis of PONV in adults undergoing surgery, diagnostic or therapeutic procedures under general or regional anesthesia or under monitored anesthetic care.

Specific objectives

- To estimate the frequency of adverse effects associated with the administration of haloperidol.
- Estimate the need for therapeutic antiemetic agents following the prophylactic administration of haloperidol for PONV.
- To assess whether the risk of PONV changes depending on the route of administration of haloperidol.

- To evaluate whether the risk of PONV changes depending to the time of administration of haloperidol.
- To evaluate whether the risk of PONV changes according to the dose of haloperidol administered.

Materials and methods

Design

Systematic literature review and met-analysis of clinical trials.

Inclusion criteria

- Type of studies: Randomized clinical trials, both published and unpublished, evaluating the effectiveness of haloperidol in the prevention of PONV.
- Type of participants: Adult patients undergoing diagnostic or therapeutic procedures under general/regional anesthesia or sedation.
- Types of interventions: Haloperidol, at any dose or route
 of administration, any moment prior to the occurrence of
 PONV versus placebo, another drug or no treatment. The
 drug could be administered in the preoperative period, during the induction of anesthesia, in the intraoperative or
 postoperative period (prior to the occurrence of nausea
 and/or vomiting).
- Types of outcomes: Incidence of nausea, vomiting, PONV, rescue remedies requirement, Q-T segment disorders and any haloperidol-related adverse effect.

Exclusion criteria

- Studies evaluating combined therapies in the haloperidol group, without specifying individual antiemetic effects.
- Trials with less than 20 participants.

Search methods

The e-search of the literature was performed in the following databases: Medline, EMBASE, CINAHL and The Cochrane Controlled Trials Register. The search used a combination of key words (as described in the search terms) and a filter recommended in Medline's Clinical Queries section with 93% sensitivity and 97% specificity to identify randomized clinical trials. The search had no restrictions with regard to language or year of publication.

Search terms

- (MESH) AND haloperidol (MESH) or postoperative (ALL FIELDS) AND nausea (ALL FIELDS) AND vomiting (ALL FIELDS) AND Haloperidol (ALL FIELDS) MeSH-NAUSEA OR NAUSEA* OR INAPPETENCE
- 2. MeSH-VOMITING OR VOMIT* OR EMESIS OR EMET*
- 3. MeSH-POSTOPERATIVE NAUSEA AND VOMITING OR POST-OPERATIVE NAUSEA AND VOMITING
- 4. Randomized Controlled Trials
- 5. #1 OR #2 OR #3
- 6. MeSH-POSTOPERATIVE OR POST-OPERATIVE

- MeSH-ANESTHESIA OR ANAESTHESIA OR ANESTHET* OR ANAESTHET*
- 8. MeSH-HALOPERIDOL

Filter: (randomized controlled trial [Publication Type] OR (randomized[Title/Abstract AND controlled[Title/Abstract]).

A search of the unpublished literature was accomplished by contacting the most representative authors and the haloperidol pharmaceutical manufacturers. The search was done on the references of the selected articles, PONV clinical practice guidelines, editorials and relevant review articles.

Two independent researches carried out the search and any discrepancies were settled by consensus.

Selection of trials

The titles of the articles found in the search were reviewed to identify any relevant articles. Then a selection by abstract was done, and an attempt was made to get the full text of the selected article. The inclusion and exclusion criteria by two independent authors were used and any conflicts were settled by consensus.

Data extraction

Two independent researchers reviewed the selected articles in order to extract the information about the participants, the interventions and outcomes. Any differences were agreed upon by consensus.

Quality evaluation of the trials included

The quality evaluation of the selected trials was performed using the instrument suggested by the GRADE Guidelines (Guyatt G.H.) and the CONSORT 2010 Statement checklist (Murphy J.F.), for the evaluation of clinical controlled trials. Two independent reviewers carried out the evaluation and any differences were settled by consensus.

Data analysis

Initially, a descriptive analysis of the information extracted and the quality of the primary trials was performed. The quantitative analysis was done using the Cochrane collaboration RevMan5 software.

Management of missing data

An attempt was made to contact the authors in order to obtain the missing data of the primary studies. If the data could not be obtained, the information was excluded from the analysis. The selected trials for which the complete text was not available (Maggi and Dyberg) were included, despite the inability to complete the planned quality analysis; the data were obtained from the systematic review published by Buttner in 2004. The decision to include the data was made based on the fact that the sample size of the relevant references was representative for the analysis. A sensitivity analysis was done, excluding

these articles, in order to measure their impact on the overall effectiveness estimate.

Approximation to heterogeneity

Clinical

A visual exploration based on the descriptive analysis of the articles included to estimate the variability of the demographic data and the risk factors for PONV was accomplished. The adjustment of the clinically relevant variables was done through sensitivity analysis.

The measurement of the meta-analytical estimates was done using the random effects model. The heterogeneity estimate was performed using the $\rm I^2$ test.

Subgroup analysis

The subgroup analysis was performed on the basis of the outcome measurements times, the type of antiemetic control, time of administration of haloperidol, and the route of administration of haloperidol.

Sensitivity analysis

A sensitivity analysis was carried out to assess the impact of the data included from sources of articles not available in their complete text and based on the methodological quality of primary studies.

Results

395 articles were identified through our own search strategy. 38 articles were selected by title and abstract, three of which were repeated in every database, 12 more were repeated in two of the databases enquired and 5 articles were in just one database. Of the 20 original articles obtained, 7 were excluded because of combined therapy in the haloperidol group with no differential analysis; 2 were excluded due to the use of therapeutic haloperidol – not prophylactic – and another one was excluded because of failing to consider the PONV scenario. By the end of the selection process, 10 articles were obtained, and an attempt was made to obtain the complete text through the Pontificia Universidad Javeriana library.

Failure to obtain the full text led to an attempt to contact the authors. However, the full texts of the Dyberg 1962 and Maggi 1964 articles were impossible to retrieve. The frequency of the outcomes was obtained from the systematic review published by Buttner in 2004. No quality analysis of these trials was done; nevertheless, they were included in the analysis because their sample size was representative.

During the manual search, no additional studies were identified. As a result of the communication established with Jansen, the pharmaceutical company that produces Haloperidol, and with the author, Koung-Shing Chu, from the Department of Anesthesiology of Kaohsiung Medical University Hospital of Kaohsiung, Taiwan, no additional unpublished studies were identified.

Characteristics of the studies included

10 trials published since 1962 to 2010 were analyzed and these included 2711 participants. The age of the participants ranged from 15 to 75 years. One of the trials was performed in patients who had undergone laparoscopic cholecystectomy (Chu), two trials on gynecological laparoscopy (Aouad and Wang), another article in open gynecological surgery (Chu) and the rest included mixed surgical population. The doses of haloperidol used ranged from 0.25 mg to 5 mg.

The time of administration of prophylactic treatment for PONV was following the induction of anesthesia in 4 trials and during the intraoperative period in another 4 trials. Two trials did not specify the time of administration and it was impossible to obtain the information when trying to contact the authors.

The control drugs used were ondansetron, droperidol, dexametasone and placebo.

The outcomes included in the trials were incidence of nausea, vomiting, PONV, evaluated at 2h, 4h, 6h and 24h, in addition to the need for rescue antiemetic agents and prolongation of the QT interval.

Quality of the trials included

We were unable to obtain data on the quality of the experiments published by Dyberg in 1962 and Maggi 1964. The outcome data were obtained from the meta analysis published by Buttner in 2004 and the appendixes to this review, published over the Internet.^{22–25}

All the trials included are randomized clinical trials. Six of the trials included reported patients lost to follow up, but none of these losses impacted the validity of the results. Two of the trials had no patients lost to follow up and other two trials do not report any related data. Only three trials use the intention-to-treat analysis but the rest do not report any related data.^{26–29}

Eight of the trials included report that the participants were blind; eight trials report that the treating physicians were blind and in six the evaluations were blind. Six of the trials specify the development and blind nature of the randomization process. $^{26-34}$

The quality of the experiments and the amount of information reported in the articles improved according to the year of publication, probably because of the development of checklists that facilitate the job of the researchers in organizing and assembling the reports of the trials. The articles included in this systematic review are average to high quality. Eight of the articles meet 5 or more of the 7 items evaluated under Guyatt's scale. The two remaining articles (Tornetta and Maggi) do not meet the quality criteria evaluated. The quality evaluation suggested by CONSORT and applied to our meta analysis highlights the fact that 5 articles met at least 11 of the items; 1 article meets 9 of the items and 2 articles (Dyberg and Maggi) have no data available to do the quality evaluation

In view of the clinical heterogeneity accounted for by diversity of the control interventions, we decided to undertake a subgroup analysis by type of control.



Fig. 1 - Haloperidol 0.25 mg.

Haloperidol versus placebo

Six of the studies included compare haloperidol against placebo for the prophylaxis of PONV. As a whole, the reports favor the use of haloperidol, but it was not possible to do the quantitative synthesis due to the broad clinical heterogeneity resulting from the broad range of doses used; hence, we decided to do the per dose analysis.

Haloperidol 0.25 mg

Just Tornetta in 1972 evaluated the prophylactic effect of I.M. haloperidol 0.25 mg, 45 min before the induction of anesthesia (Fig. 1).

There is no statistically significant difference in the incidence of PONV in the first 6 postoperative hours, which was the only time period evaluated. However, Buttner reported that there is no significant difference in PONV at 24 h: RR 0.57 95% CI (0.18–1.28) for nausea and RR 0.85 (0.4–1.79) for vomiting.

Haloperidol 0.5 mg

Only Tornetta used I.M. haloperidol 0.5 mg, 45 minutes prior to surgery (Fig. 2).

The control group has a larger number of participants. This study was developed in two phases: the initial phase included 4 groups with doses of 0.5, 1 and 2 mg, versus placebo; the second phase evaluated 0.25 mg and 4 mg doses versus placebo.

There is a significant difference in the prevention of PONV at 6 h. According to Buttner, the differential analysis of the incidence of nausea and vomiting at 24 h indicates a RR 0.48 95 CI % (0.24–0.96) for nausea and RR 0.28 95% CI (0.08–0.63) for vomiting. There is no differential decrease in the incidence of nausea or vomiting (Fig. 3).

Aouad studied the participants who underwent laparoscopic surgery; Parlow focused on urological and lower limbs surgery under regional anesthesia with intrathecal morphine and Wang focused on gynecological laparoscopy. The total effect shows benefit with the administration of haloperidol 1 mg in the prevention of nausea and vomiting at 24 h. There is moderate heterogeneity, probably accounted for in Parlow's trial (intrathecal morphine-associated anesthesia) with a baseline risk of PONV different from the other trials included in the comparison; nevertheless, we do not know the PONV risk difference among the various types of surgeries and anesthetic techniques.

Haloperidol's route of administration in Aouad and Wang's trials is intravenous, while in Parlow's it is intramuscular, which could account for the heterogeneity. Tornetta was excluded from this analysis because he only evaluated the outcome after 6 h.

In conclusion, there is a 30% PONV risk reduction with haloperidol versus placebo 95% CI (0.3–49%, P: 0.03).

Need for rescue antiemetic agents

In Aouad's trial, patients initially received intravenous promethazine 12.5 mg and ondansetron 4 mg when symptoms persisted. In Parlow's trial, patients received I.V. diphenhydrinate 50 mg and if symptoms persisted they were administered I.V. prochloperazine 10 mg and ondansetron 4 mg. Wang's trial does not report the specific rescue antiemetic agents (Fig. 4).

Haloperidol decreases the need for rescue antiemetic agents by 60%. The major effect is evidenced in Wang's trial with a higher weight of the analysis because of a larger sample size. The uncertainty in the other trials results from the small sample size; however, the specific estimates favor the use of haloperidol. There is no heterogeneity among the trials included in this analysis

QT interval

Aouad and Wang's trials measured the QT interval and showed no increases in the measurement following the administration of haloperidol. Wang took the measurement 10 min following the drug administration, while Aouad did it at the end of surgery.

Sedation

Aouad and Wang's trials measured sedation up to 2h after surgery, using the 0–10 numerical rating scale; there were no differences in the sedation rating scores between the groups (Fig. 5).

This analysis included the trials comparing haloperidol's 2 mg effectiveness versus placebo in the prophylaxis of PONV at 24 h. The overall estimate favors the haloperidol group, with statistical significance, reduced incidence of PONV by 34% at 24 h and 95% CI (18–49%). The effect is the same for nausea 0.71 95% CI (0.51–0.99) and vomiting 0.61 95% CI (0.41–0.89) (Fig. 6).

In this analysis we included the mean outcomes at $2\,h$ by Chu, at $12\,h$ by Parlow and at $6\,h$ by Tornetta. In the light of the heterogeneity, it was not possible to do a meta-analytical summary; however, two of the trials do evidence the protective effect of haloperidol at $6\,and\,12\,h$. There is no statistically

	Haloperidol		Placebo			Risk ratio	Risk ratio	
Study o subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI	
Tornetta 1972	9	50	22	48	100.0%	0.39 [0.20, 0.77]		

Fig. 2 - Haloperidol 0.5 mg.

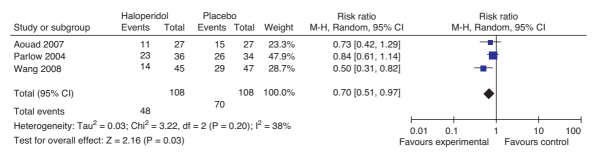


Fig. 3 - Haloperidol 1 mg.

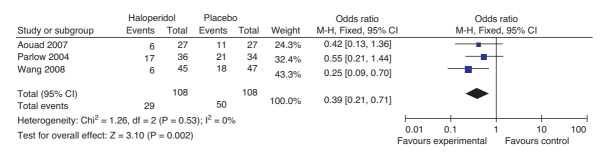


Fig. 4 - Need of rescue antiemetics.



Fig. 5 - Haloperidol 2 mg.

	Haloperidol		Placebo			Risk ratio	Risk ratio	
Study or subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI	
Chu 2008	5	72	9	75	23.1%	0.58 [0.20, 1.64]		
Parlow 2004	19	38	26	34	47.0%	0.65 [0.45, 0.95]	-	
Tornetta 1972	6	52	22	48	29.9%	0.25 [0.11, 057]		

Fig. 6 - Haloperidol 2 mg vs. placebo in the first 12 h.

significant difference in the incidence of PONV in the first 2 h (Fig. 7).

Chu used ondansetron 4 mg as rescue medication, while Parlow used I.V. diphenhydrinate 50 mg; when symptoms persisted, I.V. prochloperazine 10 mg and ondansetrón 4 mg were administered. There is evidence of a reduced need for the use of rescue agents in the patients receiving haloperidol versus placebo: 53% CI (18–73%).

QT interval

Chu measured the QTc interval before and 10 min after the administration of haloperidol. No changes were found in the length of the interval following the administration of the agents.

Haloperidol 4 mg

The studies comparing haloperidol 4 mg against placebo are Tornetta 1972 and Maggi 1964. The data are taken from Buttner's 2004 meta-analysis (Fig. 8).

Maggi randomized 140 patients to receive either haloperidol 4 mg or placebo at the end of surgery. Tornetta administered 4 mg of I.M. haloperidol prior to the induction of anesthesia. A 38% reduction in the incidence of postoperative nausea and vomiting was observed 95% CI (36–78%). Despite the difference in the routes of administration, the effect is maintained.

Haloperidol 5 mg

Dyberg evaluated the effectiveness of I.V. haloperidol 5 mg, administered during the course of surgery for prophylaxis

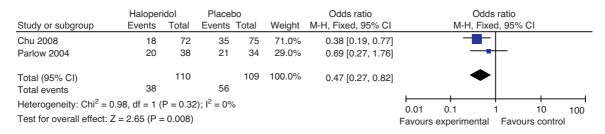


Fig. 7 - Need for rescue antiemetic agents.

Study or subgroup	Halop Events	eridol Total	Placeb Events		Weight	Risk ratio M-H, random, 95% CI	Risk ratio M-H, random, 9	-
Maggi 1964	11	70	25	70	72.8%	0.44 [0.24, 0.82]	-	
Tornetta 1972	4	51	16	51	27.2%	0.25 [0.09, 0.70]		
Total (95% CI)		121		121	100.0%	0.38 [0.22, 0.64]	•	
Total events	15		41					
Heterogeneity: Tau ²	= 0.00; Chi ²	= 0.86, 0	df = 1 (P =	= 0.35)		0.01 0.1 1	10 100	
Test for overall effec	t: Z = 3.57 (F	o = 0.000)4)		F	avours experimental F	avours control	

Fig. 8 - Haloperidol 4 mg.

of PONV in a mixed surgical population. Follow up was for 24 h. The data are taken from Buttner's systematic review and show a reduction in the risk of nausea of 63% 95% CI (54–70% P < 0.00001) and a 70% reduction CI (59–78% P < 0.00001).

Haloperidol versus droperidol

Chu and Wang compared the effectiveness of haloperidol and droperidol for the prophylaxis of PONV. Chu included women over 18 years of age, undergoing laparoscopy-assisted vaginal hysterectomy to compare I.V. haloperidol 2 mg versus droperidol 1.25 mg, 15 min after the induction of anesthesia. Wang randomized adult women undergoing gynecological laparoscopy to either I.V. haloperidol 1 mg or droperidol 0.625, 15 min prior to the induction of anesthesia (Fig. 9).

There is no difference in the incidence of PONV at $4\,h$, RR 0.97 95% CI 0.52–1.79 P: 0.91, more between $4\,and\,24\,h$ RR 0.99 95% CI (0.64–1.53 P: 0.96).

Need for rescue agents

I.V. Ondansetrón 4 mg was administered in both studies as rescue medicine to patients reporting intolerable nausea and vomiting (Fig. 10).

There is no statistically significant difference in the need for rescue medicines between groups in both trials. The measurements for this outcome were taken at 4 h in Wong's trial and at 24 in Chu's.

QTc interval

There are no reports of QTc interval prolongation or length, before and after the administration of the antiemetic agents.

Haloperidol contra ondansetrón

Incidence of PONV at 24 h

Three studies compared the effectiveness of haloperidol and ondansetron for the prophylaxis of PONV. Aouad randomized adult women undergoing gynecological surgery to receive I.V. haloperidol 1 mg or ondansetron 4 mg, 10 min after the induction of anesthesia. Feng randomized men and women undergoing laparoscopic cholecystectomy to receive haloperidol 2 mg, 30 min before the end of surgery or ondasentron 4 mg, following the induction of anesthesia. Lee randomized the mixed surgical population under general anesthesia to receive I.V. haloperidol 2 mg or ondansetrón 4 mg, 30 min before the end of surgery (Fig. 11).

There is no statistically significant difference in the incidence of PONV at 24 h, in the incidence of nausea RR 0.86 (0.57–1.29 P: 0.46), or the incidence of vomiting RR 1.24 95% CI (0.66–2.35) P: 0.5, at 24 h.

Rescue medicines

In Aouad's trial, patients received I.V. promethazine 12.5 mg as rescue agent when the score in the nausea rating scale exceeded 2/10 for over 10 min. If the symptoms persisted for over 10 min, an additional dose of ondansetron 4 mg was administered. In Lee's trial, I.V. 25 mg metochlopramide were administered as rescue agent when an episode of moderate to severe nausea or severe vomiting occurred. In Feng's trial I.M. 10 mg metochlopramide were administered when patients presented with nausea scores above 5/10, if the patient vomited or at the request of the patient (Fig. 12).

There is no difference in the need for rescue medicines at $24\,\mathrm{h.}$

QT interval

Aouad and Lee did not find any participant with a QTc interval over 470 ms or cardiac dysrrhythmia. No changes were

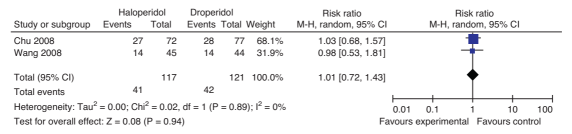


Fig. 9 - Incidence of PONV at 24 h.

Study or subgroup	Haloperi Events	dol Total	Droperidol Events Total Weight			Risk ratio M-H, random, 95% CI	Risk ratio I M-H, random, 95% CI
Chu 2008	18	72	20	77	77.1%	0.96 [0.56, 1.67]	-
Wang 2008	6	45	7	44	22.9%	0.84 [0.31, 2.30]	· -
Total (95% CI)		117		121	100.0%	0.93 [0.58, 1.51]	•
Total events	24		27				
Heterogeneity: Tau ² = Test for overall effect:			= 1 (P = 0	.81); I	F	0.01 0.1 1 10 100 Favours experimental Favours control	

Fig. 10 - Need for rescue medicines.



Fig. 11 - Incidence of PONV at 24 h.

Objects and an explanation	Haloperidol		Ondansetron			Odds ratio	Odds ratio I M-H, random, 95% C		
Study or subgroup	Events	Total	Events	Total	Weight	M-H, random, 95% CI	IVI-H, ran	30m, 95% CI	
Aouad 2007	6	27	12	27	27.3%	0.36 [0.11, 1.17]		+	
Feng 2009	17	64	14	65	41.0%	1.32 [0.59, 2.96]	_	+	
Lee 2007	10	43	8	43	31.7%	1.33 [0.47, 3.77]	_	 	
Total (95% CI)		134		135	100.0%	0.92 [0.42, 2.01]	<		
Total events	33		34						
Heterogeneity: Tau ² =	0.21; Chi ² =	3.63, df =	2 (P = 0.16	0.0	01 0.1	1 10	100		
Test for overall effect:	Z = 0.20 (P =	= 0.84)		***	Favours experimental Favour				

Fig. 12 - Rescue medicines.

detected in the length of the QTc interval, before or after the administration of the antiemetic agents.

Extrapyramidalism

None of the patients in these three studies presented extrapyramidal side effects.

Time of administration of haloperidol

Yang in 2008 had the goal to measure the impact of time of administration of haloperidol on its effectiveness as antiemetic prophylaxis. 94 patients between 20 and 65 years

of age participated in the study; these patients had been scheduled for gynecological surgery, thyroid, breast or plastic surgery. One group received I.V. haloperidol 2 mg at the start of surgery and placebo 2 ml at the end of the procedure. The second group received 2 ml of I.V. placebo at the start of surgery and I.V. haloperidol 2 mg at the end of the procedure. No statistically significant differences were found with regard to the incidence of nausea, vomiting, PONV or need for rescue antiemetic agents at 2 h and between 2 and 24 h. Neither was there any difference in the sedation scores at 2 h.

Discussion

Ten clinical controlled trials including 2711 patients randomized to haloperidol or control therapy were identified. Emphasis was placed on identifying unpublished trials and no articles were excluded based on year of publication, language in which the trial was reported or search strategy used. The manual search for references, the contact with the pharmaceutical industry and with recognized authors on the topic did not result in non-identified articles in the e-search strategy.

Haloperidol's effectiveness as prophylaxis for PONV is supported by this systematic review, and no statistically significant differences were found when comparing ondansetron versus droperidol.

In the light of the available evidence, we were unable to determine the differential effect of haloperidol based on the duration and type of surgery and the type of anesthesia. The clinical heterogenity found in the study can be explained because of the baseline risk was variable in the participants of the trials (gender, history of smoking, history of PONV or kinetosis, use of opiates in the perioperative period), because of the different doses administered, the time of administration, routes of administration, type of surgery, duration of the procedure and type of anesthesia. However, this surgical population, diverse in terms the relevant characteristics for the outcome, may increase the overall results in a uniform and consistent manner in favor of haloperidol.

No primary trials in which haloperidol was combined with different modes of action (combined prophylaxis) in the same group were included, without evaluating the effect in a differential manner for the same group. However, in moderate to high-risk patients for PONV, the combination of 2 or 3+ mechanisms of action may enhance the postoperative outcomes. These results may not be extrapolated to patients requiring combined prophylaxis – a group of people that will probably benefit further with the prophylactic use of haloperidol. 35–37

The availability and the cost of serotoninergic antagonists such as ond ansetron have improved in last few years in the various countries (generic alternatives less than one dollar or its equivalent). $^{38-40}$

Conclusion

Haloperidol's beneficial effect for the prevention of PONV in the first 24 h post-op is absolutely evident. Despite the use of different doses administered and different routes of administration uses in the trials included, the results are consistent in favor of effectiveness. The minimum effective dose is 0.5 mg and the effectiveness is incremental with respect to the dose. This increase can be better appreciated when raising the dose from 2 mg to 4 mg; additional trials are hence required in order to establish the safety of these doses. However, doses of 1–2 mg are a safe and effective alternative for the prophylaxis of PONV. The dose of haloperidol used for the prophylaxis of PONV is lower than the dose used in the treatment of psychiatric diseases – an area in which the agent is widely used chronically. Consequently, smaller and single doses should not be associated with serious complications such as arrhythmias, hence

increasing the risks and costs in the post-anesthesia care unit, during the postoperative hospital stay or within the outpatient environment.

Additionally, when comparing against all the available antiemetic agents such as dexamethasone and ondansetron, haloperidol represents an effective and safe alternative for the prophylaxis of NOGP. The cost and extended half-life of the product could make it more attractive in the perioperative environment. Cost-effectiveness trials are required in our environment, to be able to draw final conclusions.

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Conflict of interest

None of the authors reported any conflicts of interests. All the authors perform health-care and academic activities unrelated to the production or sales of drugs.

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